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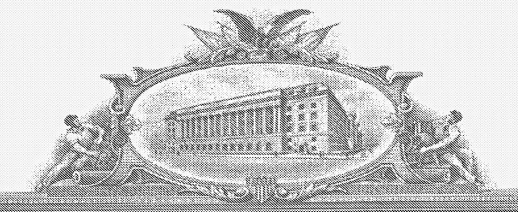
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Respectfully submitted,

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Date: June 27, 2003

REGISTRATION NO. 47,042

(if appropriate)

#### TASTE RECEPTOR OF THE T1R FAMILY FROM DOMESTIC CAT

#### FIELD OF THE INVENTION

[0001] The present invention relates to the field of sensory mechanisms of the domestic cat, Felis catus. The invention relates, for example, to the discovery of a gene of Felis catus encoding a taste receptor of the T1R family, specifically T1R3. The invention further relates to the polypeptide encoded by the T1R3 gene of the domestic cat, and to methods and uses of the same.

### **BACKGROUND OF THE INVENTION**

[0002] The sense of taste is important for determining food choice, for regulating food intake, and for ensuring efficient use of ingested nutrients. Taste can act as a warning system for the

presence of potentially harmful foods, by, for example, the aversive sensations of sourness or bitterness, and as an attractant to potentially nutrient-rich foods, by, for example, the appealing sensations of sweetness, saltiness, and umami.

[0003] Taste stimuli are received by taste receptor cells assembled into taste buds that are located in the epithelium of taste papillae of the tongue (Kitagawa et al., Bioch. Bioph. Res. Comm., 283:236-242 (2001)). The stimuli are believed to be transduced by taste receptors at the surface of the taste receptor cells (Id.). The taste receptors encoded by the genes of a given species are reflective of that species' food choices. For example, the "sweet receptors" of an herbivorous species are expected to be different from those of a carnivorous species, since the two consume completely different diets whose foods contain different primary stimuli. Since taste receptor specificity likely reflects food choice, it follows that receptor sequence homology among species may be as predictive or more predictive of food preferences of a given species as phylogenetic relatedness among species.

[0004] The behavior of the domestic cat (*Felis catus*), a carnivore, towards stimuli such as sweet carbohydrates, which it generally cannot taste, and towards L-amino acids, which it generally can taste, should be explicable by the specificity of taste receptors of other carnivores. Direct knowledge of taste receptor genes will allow insight into an animal's sensory world and may be useful for identifying modulators of the taste receptors encoded thereby to influence an animal's taste preferences.

[0005] Molecular receptors for the taste element of sweetness have recently been identified from human, mouse, and rat. Thus far, there are three known members of the T1R taste receptor family: T1R1, T1R2, and T1R3 (Montmayeur & Matsunami, Curr. Opin. Neurobiol., 12:366-371 (2002)). The T1Rs are G-protein coupled receptors with long N-terminal extracellular domains believed to be involved in ligand binding (Id.). The T1R3 receptor is located within the Sac locus, the primary genetic locus controlling preference for sweet-tasting stimuli in mice (Li et al., Mamm. Genome, 12:13-16 (2001); Li et al., Mamm. Genome, 13:5-19 (2002)). Within the cell, the taste receptors heterodimerize, with T1R3 being required for the activity of T1R1 and T1R2. In mouse, the T1R1/T1R3 heterodimer functions as a receptor for selected amino acids, while the T1R2/T1R3 heterodimer functions as a receptor for stimuli considered sweet by humans. It appears that the T1R3 component couples the taste receptor to cellular signal transduction processes, thereby ensuring that the stimulus-binding event is transduced to a neural signal. Thus, knowledge of the T1R3 receptor may lead to better understanding of species-specific reactions to sapid stimuli.

[0006] Currently, mechanisms for identifying novel taste stimuli for the domestic cat are limited, for example, to exhaustive and difficult feeding studies in which a novel ingredient is paired with a control ingredient and intake of the two are compared. Considerable time, effort, and expense can be expended in the discovery of a single stimulus. Furthermore, feline illnesses often are exacerbated by a cat's refusal to eat. Additionally, the molecular features that define acceptable taste stimuli for domestic cat remain largely unknown, making rational computational design approaches for taste stimuli difficult. As a result, knowledge of the feline taste receptor and its ligands may lead to a better understanding of cat taste perception and modulation thereof.

[0007] The present invention provides, for example, a novel feline taste receptor, T1R3, and methods of use thereof to identify taste stimuli preferred by the cat. The screening methods of the invention allow the rapid screening of binding partners and/or modulators, thereby presenting significant cost savings. The results of the feline T1R3 receptor studies reflect the taste profile of the domestic cat.

#### SUMMARY OF THE INVENTION

[8000] Certain embodiments of the present invention relate to polynucleotides encoding a T1R3 receptor, including, but not limited to polynucleotides having the nucleotide sequence of SEQ ID NO:1, fragments of the polynucleotide of SEQ ID NO:1 encoding a polypeptide having substantially the same biological activity as a polypeptide encoded by the nucleotide sequence of SEO ID NO:1, variants of the polynucleotide of SEQ ID NO:1 having at least 80% homology to the polynucleotide of SEQ ID NO:1 and encoding a polypeptide having substantially the same biological activity as a polypeptide encoded by the nucleotide sequence of SEQ ID NO:1, variants of the polynucleotide of SEQ ID NO:1 having at least 80% homology to the polynucleotide of SEQ ID NO:1 and encoding a polypeptide conferring modified taste perception to one or more taste stimuli relative to a polypeptide encoded by the polynucleotide of SEQ ID NO:1, nucleotide sequences encoding the amino acid sequence of SEQ ID NO:2, nucleotide sequences substantially complementary to the nucleotide sequence of SEQ ID NO:1, and nucleotide sequences that hybridize to the complement of the polynucleotide having SEQ ID NO:1 under high stringency conditions. The polynucleotides of the invention may be DNA or RNA and may be single- or double-stranded. In some embodiments of the invention, the polynucleotide fragments have at least about 42 nucleotides. The polynucleotide fragments of the invention encode, for example, an extracellular domain of the polypeptide of SEQ ID NO:2, a transmembrane domain of the polypeptide of SEQ ID NO:2, or an intracellular domain of the polypeptide of SEQ ID NO:2. In other embodiments of the invention, the polynucleotide variants of the polynucleotide of SEQ ID NO:1 encoding an amino acid sequence of SEQ ID NO:2 having a nonconserved amino acid substitution, for example, at residue 59 and/or residue 64.

[0009] The invention also encompasses expression vectors containing the polynucleotides of the invention operably linked to a promoter. Another embodiment of the invention provides host cells containing the expression vector. The host cells may be mammalian, including feline. The invention further encompasses cell cultures of the host cells. The invention also encompasses methods of producing feline T1R3 receptor by culturing the host cells and recovering receptor therefrom.

[0010] Another embodiment of the invention includes T1R3 receptor polypeptides encoded by polynucleotides of the invention. The polypeptides of the invention include, for example, those having the amino acid sequence of SEQ ID NO:2, fragments of at least 30 contiguous amino acids of SEQ ID NO:2, and variants thereof having substantially the same biological activity as the polypeptide of SEQ ID NO:2. The variant polypeptides of the invention may have an amino acid sequence having at least one sequence variation of SEQ ID NO:2 that confers modified taste perception to one or more taste stimuli relative to a polypeptide of SEQ ID NO:2. The invention provides methods of identifying a feline T1R3 receptor variant that confers modified taste perception by expressing a variant of the polynucleotide of SEQ ID NO:1 having at least 80% homology to the polynucleotide of SEQ ID NO:1 and detecting an increase or a decrease in the biological activity of the polypeptide encoded by the variant relative to the biological activity of the polypeptide encoded by SEQ ID NO:1.

[0011] The invention further provides kits for the detection of polynucleotides encoding a feline T1R3 receptor including a polynucleotide that specifically hybridizes to a polynucleotide encoding a polypeptide having an amino acid sequence of SEQ ID NO:2 and instructions relating to detection thereof.

[0012] Also provided by the invention are antibodies that immunoreact specifically with at least one epitope of a polypeptide of the invention. The invention also includes kits for the detection of polypeptides encoding a feline T1R3 receptor including antibodies of the invention and instructions relating to detection.

[0013] Further provided by the invention are methods for identifying compounds that interact with a feline T1R3 receptor by contacting a feline T1R3 receptor with a test compound, and

detecting interaction between the receptor and the compound. The receptor may be bound to a solid support. In one aspect of the invention, the recognition sites of the receptor are coupled with a monitoring system, either electrical or optical. In another embodiment, the solid support is formulated into a feline-specific electronic tongue.

[0014] The invention also provides methods for identifying agonists and antagonists of feline T1R3 receptor. For example, the methods of the invention include identification of an agonist of a feline T1R3 receptor by expressing an expression vector of the invention in the presence of a test compound, and detecting an increase in biological activity of a polypeptide produced by the expression in the presence of the compound relative to biological activity of the polypeptide in the absence of the compound. Also included are methods for identifying agonists of a feline T1R3 receptor by contacting a polypeptide of the invention with a test compound, and detecting an increase in biological activity of the polypeptide in the presence of the compound relative to biological activity of the polypeptide in the absence of the compound.

[0015] Methods for identifying antagonists of the polypeptides of the invention also are provided. For example, the invention provides methods for identifying antagonists of a feline T1R3 receptor by expressing an expression vector in the presence of a test compound, and detecting a decrease in biological activity of a polypeptide produced by the expression step in the presence of the compound relative to biological activity of the polypeptide in the absence of the compound. Another example of methods for identifying an antagonist of a feline T1R3 receptor involves contacting a polypeptide of the invention with a test compound, and detecting a decrease in biological activity of the polypeptide in the presence of the compound relative to biological activity of the polypeptide in the absence of the compound.

[0016] Another embodiment of the invention includes compounds and compositions for modifying the taste perception of a mammal, such as a cat. The compounds and compositions may contain at least one of polynucleotides of the invention, polypeptides of the invention, or compounds identified by the methods of the invention. Examples of the compositions of the invention include veterinary foods and drinks and pharmaceutical compositions. The compositions of the invention may include a pharmaceutically acceptable excipient. The compositions of the invention may be breed-specific. Methods for modifying the taste perception of a mammal (e.g., a cat) by administering to the mammal a polynucleotide of the invention, a polypeptide of the invention, and/or a compound identified according to the methods of the invention also are provided.

[0017] The invention further provides transgenic animals comprising a polynucleotide of the invention.

[0018] The materials, methods, and examples provided herein are illustrative only and are not intended to be limiting. Other features and advantages of the invention will be apparent from the following detailed description and claims.

#### BRIEF DESCRIPTION OF THE DRAWINGS

[0019] Figure 1 shows the multiple sequence alignment of the T1R3 receptor of domestic cat (SEQ ID NO:1) with known nucleotide sequences of receptors of the T1R family from human (T1R1, SEQ ID NO:8; T1R2, SEQ ID NO:5; T1R3, SEQ ID NO:11), mouse (T1R1, SEQ ID NO:6; T1R2, SEQ ID NO:3; T1R3, SEQ ID NO:9), and rat (T1R1, SEQ ID NO:7; T1R2, SEQ ID NO:4; T1R3, SEQ ID NO:10). An asterisk (\*) indicates a conserved nucleotide position among the sequences.

[0020] Figure 2 shows the deduced amino acid sequence of the domestic cat taste receptor, T1R3 (SEO ID NO:2), aligned with the amino acid sequences of members of the T1R receptor family from human (T1R1, SEQ ID NO:17; T1R2, SEQ ID NO:20; T1R3, SEQ ID NO:12), rat (T1R1, SEO ID NO:16; T1R2, SEO ID NO:19; T1R3, SEO ID NO:14), and mouse (T1R1, SEQ ID NO:15; T1R2, SEQ ID NO:18; T1R3, SEQ ID NO:13). An asterisk (\*) indicates a conserved nucleotide position among the sequences. A colon (:) indicates an observed conserved amino acid substitution. A period (.) indicates an observed semi-conserved amino acid substitution. The deduced amino acid sequence for cat T1R3 (SEQ ID NO:2) contains four additional amino acids at positions 11-14 relative to the homologous T1R3 receptors of mouse (SEQ ID NO:13), human (SEQ ID NO:12), and rat (SEQ ID NO:14). The deduced sequence for cat reveals a threonine in position 64, a position equivalent to amino acid 60 in mouse, and a leucine at position 59, a position equivalent to position 55 in mouse. In mouse, amino acid substitutions of a threonine at position 60 and an alanine at position 55, both positions located within the putative extracellular N-terminal domain of the polypeptide, are present in strains of mice demonstrating low preference for the sweet stimulus saccharin (Bachmanov et al., Chem. Senses, 26:925-933 (2001)). Leucine is a conservative substitution for alanine. Accordingly, the amino acid sequence differences of cat and mouse T1R3 receptor may account for functional differences that lead to different taste preferences between the two species.

[0021] Figure 3 illustrates a phylogenetic tree showing the relatedness of the domestic cat T1R3 receptor to the T1R family of receptors including human, rat, and mouse T1R1, T1R2, and T1R3. The T1R receptors of the rat and mouse are closely related, while the T1R3 receptor of human and cat diverge from rat and mouse. Interestingly, the sweet stimuli to which the rat and mouse respond are very similar, whereas those that stimulate the human and those that stimulate the cat differ from one another and from those for rat and mouse.

[0022] Figure 4 illustrates the predicted conformation of cat T1R3 receptor. The cat T1R3 receptor is a seven transmembrane receptor similar in structure to other known members of the T1R family of receptors. The structure of the feline T1R3 receptor was generated through use of a protein modeling program available at <www.ebi.ac.uk/~moeller/transmembrane.html>.

#### DETAILED DESCRIPTION OF ILLUSTRATIVE EMBODIMENTS

[0023] The reference works, patents, patent applications, and scientific literature that are referred to herein reflect in part the knowledge of those with skill in the art and are hereby incorporated by reference in their entirety to the same extent as if each was specifically and individually indicated to be incorporated by reference. Any conflict between any reference cited herein and the specific teachings of this specification shall be resolved in favor of the latter. Likewise, any conflict between an art-understood definition of a word or phrase and a definition of the word or phrase as specifically taught in this specification shall be resolved in favor of the latter.

[0024] Standard reference works setting forth the general principles of recombinant DNA technology are known to those of skill in the art (Ausubel et al., CURRENT PROTOCOLS IN MOLECULAR BIOLOGY, John Wiley & Sons, New York, 1998; Sambrook et al., MOLECULAR CLONING: A LABORATORY MANUAL, 2D ED., Cold Spring Harbor Laboratory Press, Plainview, New York, 1989; Kaufman et al., Eds., HANDBOOK OF MOLECULAR AND CELLULAR METHODS IN BIOLOGY AND MEDICINE, CRC Press, Boca Raton, 1995; McPherson, Ed., DIRECTED MUTAGENESIS: A PRACTICAL APPROACH, IRL Press, Oxford, 1991).

[0025] As used herein, "taste perception" refers to a response (e.g., biochemical, behavioral) or sensitivity of a T1R3 receptor of the invention to a taste stimulus. "Taste stimulus" as used herein refers to any compound that elicits, for example at the biochemical level (e.g., activation or inhibition of a taste receptor) or behavioral level, a taste response which would be perceived by a mammal as at least one of the five taste elements, including sweet, salty, sour, bitter, and

umami. "Taste perception" or "taste stimulus," or variants thereof, does not require, though it does include, transmission of a neural signal resulting in *in vivo* sensation of taste by a mammal. Modification of taste perception includes an alteration of (enhancement of, reduction to, or change to) a biochemical response, an ingestive response, a taste preference, or general behavior of a mammal in response to a compound.

[0026] As used herein "polynucleotide" refers to a nucleic acid molecule and includes genomic DNA, cDNA, RNA, mRNA, mixed polymers, recombinant nucleic acids, fragments and variants thereof, and the like. Polynucleotide fragments of the invention comprise at least 10, and preferably at least 12, 14, 16, 18, 20, 25, 30, 35, 40, 45, 50, 75, or 100 consecutive nucleotides of a reference polynucleotide. The polynucleotides of the invention include sense and antisense strands. The polynucleotides of the invention may be naturally occurring or non-naturally A "synthesized polynucleotide" as used herein refers to occurring polynucleotides. polynucleotides produced by purely chemical, as opposed to enzymatic, methods. "Wholly" synthesized DNA sequences are therefore produced entirely by chemical means, and "partially" synthesized DNAs embrace those wherein only portions of the resulting DNA were produced by chemical means. The polynucleotides of the invention may be single- or double-stranded. The polynucleotides of the invention may be chemically modified and may contain non-natural or derivatized nucleotide bases as will be readily appreciated by those skilled in the art. Such modifications include, for example, labels, methylation, substitution of one or more nucleotides with an analog, internucleotide modifications such as uncharged linkages (e.g., methyl phosphonates, phosphotriesters, phosphoramidates, carbamates, etc.), charged linkages (e.g., phosphorothioates, phosphorodithioates, etc.), pendent moieties (e.g., polypeptides, etc.), intercalators (e.g., acridine, psoralen, etc.), chelators, alkylators, and modified linkages (e.g., alpha anomeric nucleic acids, etc.). Also included are synthetic molecules that mimic polynucleotides in their ability to bind to a designated sequence via hydrogen bonding and other chemical interactions. Such molecules are known in the art and include, for example, those in which peptide linkages substitute for phosphate linkages in the backbone of the molecule.

[0027] "Recombinant nucleic acid" is a nucleic acid generated by combination of two segments of nucleotide sequence. The combination may be, for example, by chemical means or by genetic engineering.

[0028] As used herein, "polynucleotide amplification" refers to a broad range of techniques for increasing the number of copies of specific polynucleotide sequences. Typically, amplification of either or both strand(s) of the target nucleic acid comprises the use of one or more nucleic

acid-modifying enzymes, such as a DNA polymerase, ligase, RNA polymerase, or RNA-dependent reverse transcriptase. Examples of polynucleotide amplification include, but are not limited to, polymerase chain reaction (PCR), nucleic acid sequence based amplification (NASB), self-sustained sequence replication (3SR), strand displacement activation (SDA), ligase chain reaction, Qβ replicase system, and the like. A wide variety of alternative cloning and *in vitro* amplification methodologies are well known to those skilled in the art. Examples of these techniques are found in, for example, Berger *et al.*, *Guide to Molecular Cloning Techniques*, METHODS IN ENZYMOLOGY 152, Academic Press, Inc., San Diego, CA (Berger), which is incorporated herein by reference in its entirety.

[0029] As used herein, the term "oligonucleotide" or "primer" refers to a series of linked nucleotide residues which has a sufficient number of bases to be used in a polymerase chain reaction (PCR). This short sequence is based on (or designed from) a genomic or cDNA sequence and is used to amplify, confirm, or reveal the presence of an identical, similar, or complementary DNA or RNA in a particular cell or tissue. Oligonucleotides comprise portions of a nucleic acid sequence having at least about 10 nucleotides and as many as about 50 nucleotides, often about 12 or 15 to about 30 nucleotides. They are chemically synthesized and may be used as probes. "Primer pair" refers to a set of primers including a 5' upstream primer that hybridizes with the 5' end of a target sequence to be amplified and a 3' downstream primer that hybridizes with the complement of the 3' end of the target sequence to be amplified.

[0030] As used herein, the term "probe" refers to nucleic acid sequences of variable length, for example between at least about 10 and as many as about 6,000 nucleotides, depending on use. Probes are used in the detection of identical, similar, or complementary target nucleic acid sequences, which target sequences may be single- or double-stranded. Longer probes are usually obtained from a natural or recombinant source, are highly specific, and are much slower to hybridize than oligomers, or shorter probes. They may be single- or double-stranded and are carefully designed to have specificity in PCR, hybridization membrane-based, or ELISA-like technologies. An "overgo probe" is a DNA probe comprising two short, overlapping DNA sequences (e.g., 10-50 nucleotides each) with a complementary overlapping region (e.g., 5-15 nucleotides) that is used in an overgo hybridization strategy. For example, an overgo probe may be two 22mers with an 8 bp complementary overlap, resulting in a 36mer overgo probe. As another example, an overgo probe may be two 24mers with an 8 bp complementary overlap, resulting in a 40mer overgo probe.

[0031] As used herein, the phrase "stringent hybridization conditions" or "stringent conditions" refers to conditions under which a probe, primer, or oligonucleotide will hybridize to its target sequence, but to a minimal number of other sequences. Stringent conditions are sequencedependent and will be different in different circumstances. Longer sequences will hybridize with specificity to their proper complements at higher temperatures. Generally, stringent conditions are selected to be about 5°C lower than the thermal melting point (T<sub>m</sub>) for the specific sequence at a defined ionic strength and pH. The T<sub>m</sub> is the temperature (under defined ionic strength, pH and nucleic acid concentration) at which 50% of the probes complementary to the target sequence hybridize to the target sequence at equilibrium. Since the target sequences are generally present in excess, at T<sub>m</sub>, 50% of the probes are hybridized to their complements at equilibrium. Stringent temperature conditions will generally include temperatures in excess of 30°C, typically in excess of 37°C, and may be in excess of 45°C. Stringent salt conditions will ordinarily be less than 1.0 M, typically less than 0.5 M, and may be less than 0.2 M. Typically, stringent conditions will be those in which the salt concentration is less than about 1.0 M sodium ion, typically about 0.01 to 1.0 M sodium ion (or other salts) at pH 7.0 to 8.3 and the temperature is at least about 30°C for short probes, primers, or oligonucleotides (e.g., 10 to 50 nucleotides) and at least about 60°C for longer probes, primers, or oligonucleotides. Stringent conditions may also be achieved with the addition of destabilizing agents, such as formamide.

[0032] As used herein "antisense oligonucleotide" refers to a nucleic acid molecule that is complementary to at least a portion of a target nucleotide sequence of interest and specifically hybridizes to the target nucleotide sequence under physiological conditions. The term "double stranded RNA" or "dsRNA" as used herein refers to a double-stranded RNA molecule capable of RNA interference, including short interfering RNA (siRNA) (see for example, Bass, *Nature*, 411, 428-429 (2001); Elbashir *et al.*, *Nature*, 411, 494-498 (2001)).

[0033] As used herein, the term "complementary" refers to Watson-Crick basepairing between nucleotide units of a nucleic acid molecule.

[0034] The term "marker gene" or "reporter gene" refers to a gene encoding a product that, when expressed, confers a phenotype at the physical, morphologic, or biochemical level on a transformed cell that is easily identifiable, either directly or indirectly, by standard techniques and includes, but is not limited to, genes encoding proteins that confer resistance to toxins or antibiotics such as ampicillin, neomycin, and methotroxate; genes encoding proteins that complement auxotrophic deficiencies; and genes encoding proteins that supply critical components not available from complex media. Examples of marker genes include green

fluorescent protein (GFP), red fluorescent protein (DsRed), alkaline phosphatase (AP), β-lactamase, chloramphenicol acetyltransferase (CAT), adenosine deaminase (ADA), aminoglycoside phosphotransferase (neor, G418r) dihydrofolate reductase (DHFR), hygromycin-B-phosphotransferase (HPH), thymidine kinase (TK), lacZ (encoding β-galactosidase), luciferase (luc), and xanthine guanine phosphoribosyltransferase (XGPRT). As with many of the standard procedures associated with the practice of the invention, skilled artisans will be aware of additional sequences that can serve the function of a marker or reporter. Thus, this list is merely meant to show examples of what can be used and is not meant to limit the invention.

[0035] As used herein, the term "promoter" refers to a regulatory element that regulates, controls, or drives expression of a nucleic acid molecule of interest and can be derived from sources such as from adenovirus, SV40, parvoviruses, vaccinia virus, cytomegalovirus, or mammalian genomic DNA. Examples of suitable promoters include, but are not limited to, CMV, MSH2, trp, lac, phage, and TRNA promoters. Suitable promoters that can be used in yeast include, but are not limited to, such constitutive promoters as 3-phosphoglycerate kinase and various other glycolytic enzyme gene promoters such as enolase or glyceraldehydes-3-phosphate dehydrogenase, or such inducible promoters as the alcohol dehydrogenase 2 promoter or metallothionine promoter. Again, as with many of the standard procedures associated with the practice of the invention, skilled artisans will be aware of additional promoters that can serve the function of directing the expression of a marker or reporter. Thus, the list is merely meant to show examples of what can be used and is not meant to limit the invention.

[0036] "Operably linked" refers to juxtaposition wherein the components are in a functional relationship. For example, a promoter is operably linked or connected to a coding sequence if it controls the transcription or expression of the sequence.

[0037] The terms "polypeptide," "peptide," and "protein" are used interchangeably herein. "Polypeptide" refers to a polymer of amino acids without referring to a specific length. Polypeptides of the invention include peptide fragments, derivatives, and fusion proteins. Peptide fragments preferably have at least about 10, 15, 20, 25, 30, 35, 40, 45, 50, 60, 70, 80, 90, or 100 amino acids. Some peptide fragments of the invention are biologically active. Biological activities include immunogenicity, ligand binding, and activity associated with the reference peptide. Immunogenic peptides and fragments of the invention generate an epitope-specific immune response, wherein "epitope" refers to an immunogenic determinant of a peptide and preferably contains at least three, five, eight, nine, ten, fifteen, twenty, thirty, forty, forty-five, or fifty amino acids. Some immunogenic peptides of the invention generate an immune response

specific to that peptide. Polypeptides of the invention include naturally occurring and non-naturally occurring peptides. The term includes modified polypeptides (wherein examples of such modifications include glycosylation, acetylation, phosphorylation, carboxylation, ubiquitination, labeling, etc.), analogs (such as non-naturally occurring amino acids, substituted linkages, etc.), and functional mimetics. A variety of methods for labeling polypeptides are well known in the art and include radioactive isotopes such as <sup>32</sup>P or <sup>35</sup>S, ligands that bind to labeled antiligands (e.g., antibodies), fluorophores, chemiluminescent agents, enzymes, and antiligands.

[0038] As used herein, the term "amino acid" denotes a molecule containing both an amino group and a carboxyl group. In some embodiments, the amino acids are  $\alpha$ -,  $\beta$ -,  $\gamma$ - or  $\delta$ -amino acids, including their stereoisomers and racemates. As used herein the term "L-amino acid" denotes an  $\alpha$ -amino acid having the L configuration around the  $\alpha$ -carbon, that is, a carboxylic acid of general formula CH(COOH)(NH2)-(side chain), having the L-configuration. The term "D-amino acid" similarly denotes a carboxylic acid of general formula CH(COOH)(NH2)-(side chain), having the D-configuration around the  $\alpha$ -carbon. Side chains of L-amino acids include naturally occurring and non-naturally occurring moieties. Non-naturally occurring (i.e., unnatural) amino acid side chains are moieties that are used in place of naturally occurring amino acid side chains in, for example, amino acid analogs. Amino acid substituents may be attached, for example, through their carbonyl groups through the oxygen or carbonyl carbon thereof, or through their amino groups, or through functionalities residing on their side chain portions.

[0039] The amino acid sequences are presented in the amino (N) to carboxy (C) direction, from left to right. The N-terminal  $\alpha$ -amino group and the C-terminal  $\beta$ -carboxy groups are not depicted in the sequence. The nucleotide sequences are presented by single strands only, in the 5' to 3' direction, from left to right. Nucleotides and amino acids are represented in the manner recommended by the IUPAC-IUB Biochemical Nomenclature Commission, or amino acids are represented by their three letters code designations.

[0040] As used herein, the term "antibody" is meant to refer to complete, intact antibodies, and Fab, Fab', F(ab)<sub>2</sub>, F<sub>v</sub>, and other fragments thereof. Complete, intact antibodies include antibodies such as polyclonal antibodies, monoclonal antibodies, chimeric antibodies, and humanized antibodies, felinized antibodies, and immunologic binding equivalents thereof. The antibodies of the invention may be labeled or unlabeled. Examples of labels of antibodies include, but are not limited to, radionuclides, enzymes, substrates, cofactors, inhibitors,

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fluorescent agents, chemiluminescent agents, magnetic particles, and the like. Recombinant immunoglobulins are included in the invention.

[0041] As used herein, the term "binding" means the physical or chemical interaction between two proteins or compounds or associated proteins or compounds or combinations thereof. Binding includes ionic, non-ionic, Hydrogen bonds, Van der Waals, hydrophobic interactions, etc. The physical interaction, the binding, can be either direct or indirect, indirect being through or due to the effects of another protein or compound. Direct binding refers to interactions that do not take place through or due to the effect of another protein or compound but instead are without other substantial chemical intermediates. Binding may be detected in many different manners. As a non-limiting example, the physical binding interaction between two molecules can be detected using a labeled compound. Other methods of detecting binding are well-known to those of skill in the art.

[0042] As used herein, the term "contacting" means bringing together, either directly or indirectly, a compound into physical proximity to a molecule of interest. Contacting may occur, for example, in any number of buffers, salts, solutions, or in a cell or cell extract.

As used herein, the terms "modulates" or "modifies" means an increase or decrease in [0043] the amount, quality, or effect of a particular activity or protein. "Modulators" refer to any inhibitory or activating molecules identified using in vitro and in vivo assays for, e.g., agonists, antagonists, and their homologs, including fragments, variants, and mimetics, as defined herein, that exert substantially the same biological activity as the molecule. "Inhibitors" or "antagonists" are modulating compounds that reduce, decrease, block, prevent, delay activation, inactivate, desensitize, or downregulate the biological activity or expression of a molecule or pathway of interest. "Inducers," "activators," or "agonists" are modulating compounds that increase, induce, stimulate, open, activate, facilitate, enhance activation, sensitize, or upregulate a molecule or pathway of interest. In some preferred embodiments of the invention, the level of inhibition or upregulation of the expression or biological activity of a molecule or pathway of interest refers to a decrease (inhibition or downregulation) or increase (upregulation) of greater than about 50%, 60%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, or 99%. The inhibition or upregulation may be direct, i.e., operate on the molecule or pathway of interest itself, or indirect, i.e., operate on a molecule or pathway that affects the molecule or pathway of interest.

[0044] A "substantially pure" polynucleotide or polypeptide is substantially separated from other cellular components that naturally accompany a native (or wild-type) nucleic acid or polypeptide and/or from other impurities (e.g., agarose gel). A substantially pure protein will comprise about 60% to more than 99% w/w of a protein sample, and may be about 90%, about 95%, or about 98% pure. As used herein, the term "isolated" refers to a molecule that has been removed from its native environment. Examples of isolated nucleic acid molecules include, but are not limited to, recombinant DNA molecules contained in a vector, recombinant DNA molecules maintained in a heterologous host cell, partially or substantially purified nucleic acid molecules, and synthetic DNA or RNA molecules.

[0045] "About" as used herein refers to +/- 10% of the reference value.

As used herein, "variant" nucleotide or amino acid sequences refer to homologs, [0046] including, for example, isoforms, species variants, allelic variants, and fragments of the sequence of interest. "Homologous nucleotide sequence" or "homologous amino acid sequence," or variations thereof, refers to sequences characterized by a homology, at the nucleotide level or amino acid level, of at least about 60%, at least about 70%, at least about 75%, at least about 80%, at least about 81%, at least about 82%, at least about 83%, at least about 84%, at least about 85%, preferably at least about 90%, at least about 95%, at least about 98%, or at least about 99%, and more preferably 100%, to a reference sequence, or portion or fragment thereof encoding or having a functional domain, including, for example, but not limited to the nucleic acid sequence of SEQ ID NO:1, or a portion of SEQ ID NO:1, which encodes a functional domain of the encoded polypeptide, SEQ ID NO:2, or to the polypeptide having amino acid sequence SEQ ID NO:2, or fragments thereof having functional domains of the full-length polypeptide. Examples of functional domains of the T1R3 polypeptide of SEQ ID NO:2 include the extracellular domains (residues 1-571, 628-641, 705-730, and 787-794 of SEQ ID NO:2), the transmembrane domains (residues 572-594, 610-627, 642-664, 681-704, 731-754, 767-780, and 795-812 of SEO ID NO:2), and the intracellular domains (residues 595-609, 665-680, 755-766, and 813-865 of SEO ID NO:2). Isoforms can be expressed in different tissues of the same organism as a result of, for example, alternative splicing of RNA. Alternatively, isoforms can be encoded by different genes. Homologous nucleotide sequences include nucleotide sequences encoding for a species variant of a protein. Homologous nucleotide sequences also include, but are not limited to, naturally occurring allelic variations and mutations of the nucleotide sequences set forth herein. Study of mutations and polymorphisms of the T1R3 receptor polynucleotide sequence may explain breed-specific and/or individual taste preferences of a mammal such as a cat. Additionally, sequence variants of the T1R3 receptor may be associated with specific disease states, such that knowledge of the gene allows diagnosis and treatment of T1R3-associated disorders (e.g., obesity, diabetes). Homologous amino acid sequences include those amino acid sequences which encode conservative amino acid substitutions in polypeptides having amino acid sequence of SEQ ID NO:2, as well as in polypeptides identified according to the methods of the invention. Percent homology may be determined by, for example, the Gap program (Wisconsin Sequence Analysis Package, Version 8 for Unix, Genetics Computer Group, University Research Park, Madison Wis.), using the default settings, which uses the algorithm of Smith and Waterman (Smith and Waterman, Adv. Appl. Math., 2: 482-489, 1981). Nucleic acid fragments of the invention preferably have at least about 5, at least about 10, at least about 15, at least about 20, at least about 25, at least about 50, or at least about 100 nucleotides of the reference nucleotide sequence. The nucleic acid fragments of the invention may encode a polypeptide having at least one biological property, or function, that is substantially similar to a biological property of the polypeptide encoded by the full-length nucleic acid sequence.

[0047] As is well known in the art, because of the degeneracy of the genetic code, there are numerous DNA and RNA molecules that can code for the same polypeptide as that encoded by a nucleotide sequence of interest. The present invention, therefore, contemplates those other DNA and RNA molecules which, on expression, encode a polypeptide encoded by the nucleic acid molecule of interest. DNA and RNA molecules other than those specifically disclosed herein characterized simply by a change in a codon for a particular amino acid, are within the scope of this invention.

[0048] Amino acid "insertions", "substitutions" or "deletions" are changes to or within an amino acid sequence. The variation allowed in a particular amino acid sequence may be experimentally determined by producing the peptide synthetically or by systematically making insertions, deletions, or substitutions of nucleotides in the nucleic acid sequence using recombinant DNA techniques. Alterations of the naturally occurring amino acid sequence can be accomplished by any of a number of known techniques. For example, mutations can be introduced into the polynucleotide encoding a polypeptide at particular locations by procedures well known to the skilled artisan, such as oligonucleotide-directed mutagenesis, which is described by U.S. Pat. Nos. 4,518,584 and 4,737,462.

[0049] A polypeptide variant of the present invention may exhibit substantially the biological activity of a naturally occurring reference polypeptide. "Biological activity" as used herein refers to the level of a particular function (for example, enzymatic activity) of a molecule or pathway of

interest in a biological system. "Wild-type biological activity" refers to the normal level of function of a molecule or pathway of interest. "Reduced biological activity" refers to a decreased level of function of a molecule or pathway of interest relative to a reference level of biological activity of that molecule or pathway. For example, reduced biological activity may refer to a decreased level of biological activity relative to the wild-type biological activity of a molecule or pathway of interest. "Increased biological activity" refers to an increased level of function of a molecule or pathway of interest relative to a reference level of biological activity of that molecule or pathway. For example, increased biological activity may refer to an increased level of biological activity relative to the wild-type biological activity of a molecule or pathway of interest. Reference to exhibiting "substantially the biological activity of a naturally occurring polypeptide" indicates that variants within the scope of the invention can comprise conservatively substituted sequences, meaning that one or more amino acid residues of a polypeptide are replaced by different residues that do not alter the secondary and/or tertiary structure of the polypeptide. Such substitutions may include the replacement of an amino acid by a residue having similar physicochemical properties, such as substituting one aliphatic residue (Ile, Val, Leu or Ala) for another, or substitution between basic residues Lys and Arg, acidic residues Glu and Asp, amide residues Gln and Asn, hydroxyl residues Ser and Tyr, or aromatic residues Phe and Tyr. Further information regarding making phenotypically silent amino acid exchanges are known in the art (Bowie et al., Science, 247: 1306-1310, 1990). Other polypeptide homologs which might retain substantially the biological activities of the reference polypeptide are those where amino acid substitutions have been made in areas outside functional regions of the protein.

[0050] A nucleotide and/or amino acid sequence of a nucleic acid molecule or polypeptide employed in the invention or of a compound identified by the screening method of the invention may be used to search a nucleotide and amino acid sequence databank for regions of similarity using Gapped BLAST (Altschul et al., Nuc. Acids Res., 25: 3389, 1997). Briefly, the BLAST algorithm, which stands for Basic Local Alignment Search Tool is suitable for determining sequence similarity (Altschul et al., J Mol. Biol., 215: 403-410, 1990). Software or performing BLAST analyses is publicly available through the National Center for Biotechnology Information (http://www.ncbi.nlm.nih.gov/). This algorithm involves first identifying high scoring sequence pair (HSPs) by identifying short words of length W in the query sequence that either match or satisfy some positive-valued threshold score T when aligned with a word of the same length in a database sequence. T is referred to as the neighborhood word score threshold (Altschul et al., J Mol. Biol., 215: 403-410, 1990). These initial neighborhood word hits act as

seeds for initiating searches to find HSPs containing them. The word hits are extended in both directions along each sequence for as far as the cumulative alignment score can be increased. Extension for the word hits in each direction are halted when: 1) the cumulative alignment score falls off by the quantity X from its maximum achieved value; 2) the cumulative score goes to zero or below, due to the accumulation of one or more negative-scoring residue alignments; or 3) the end of either sequence is reached. The BLAST algorithm parameters W, T, and X determine the sensitivity and speed of the alignment. The BLAST program uses as defaults a word length (W) of 11, the BLOSUM62 scoring matrix (Henikoff et al., Proc. Natl. Acad. Sci. USA, 89: 10915-10919, 1992) alignments (B) of 50, expectation (E) of 10, M=5, N=4, and a comparison of both strands. The BLAST algorithm (Karlin et al., Proc. Natl. Acad. Sci. USA, 90: 5873-5787, 1993) and Gapped BLAST perform a statistical analysis of the similarity between two sequences. One measure of similarity provided by the BLAST algorithm is the smallest sum probability (P(N)), which provides an indication of the probability by which a match between two nucleotide or amino acid sequences would occur by chance. For example, a nucleic acid is considered similar to a gene or cDNA if the smallest sum probability in comparison of the test nucleic acid to the reference nucleic acid is less than about 1, preferably less than about 0.1, more preferably less than about 0.01, and most preferably less than about 0.001.

[0051] The term "mimetic" as used herein refers to a compound that is sterically similar to a reference compound. Mimetics are structural and functional equivalents to the reference compounds.

[0052] The terms "patient" and "subject" are used interchangeably herein and include, but are not limited to, avians, felines, canines, bovines, ovines, porcines, equines, rodents, simians, and humans. "Host cell" includes, for example, a mammalian cell (e.g., human, rodent, feline), yeast cell, or plant cell. "Rodents" include, for example, rats and mice.

[0053] The term "treatment" as used herein refers to any indicia of success of prevention, treatment, or amelioration of a disease or condition. Treatment includes any objective or subjective parameter, such as, but not limited to, abatement, remission, normalization of receptor activity, reduction in the number of infectious particles in a patient, reduction in the number or severity of symptoms or side effects, an increase in the tolerance of the patient to an infection, or slowing of the rate of degeneration or decline of the patient. Treatment also includes a prevention of the onset of symptoms in a patient that may be at increased risk of infection but does not yet experience or exhibit symptoms thereof.

[0054] As used herein, the term "compound" means any identifiable chemical or molecule, including, but not limited to a small molecule, peptide, protein, sugar, nucleotide, or nucleic acid. Such compound can be natural or synthetic.

#### **Polynucleotides**

[0055] The invention provides purified and isolated polynucleotides (e.g., cDNA, genomic DNA, synthetic DNA, RNA, or combinations thereof, whether single- or double-stranded) that comprise a nucleotide sequence encoding the amino acid sequence of the polypeptides of the invention. Such polynucleotides are useful for recombinantly expressing the receptor and also for detecting expression of the receptor in cells (e.g., using Northern hybridization and in situ hybridization assays). Such polynucleotides also are useful in the design of antisense and other molecules for the suppression of the expression of T1R3 receptor in a cultured cell, a tissue, or an animal; for therapeutic purposes; or to provide a model for diseases or conditions characterized by aberrant T1R3 expression. Specifically excluded from the definition of polynucleotides of the invention are entire isolated, non-recombinant native chromosomes of host cells. One embodiment of polynucleotides of the invention is the nucleotide sequence of SEQ ID NO:1. It will be appreciated that numerous other polynucleotide sequences exist that also encode the T1R3 receptor of the invention due to the well-known degeneracy of the universal genetic code.

[0056] The invention also provides a purified and isolated polynucleotide comprising a nucleotide sequence that encodes a mammalian (e.g., feline) polypeptide, wherein the polynucleotide hybridizes to a polynucleotide having a sequence of SEQ ID NO:1, or the non-coding strand complementary thereto, under stringent hybridization conditions.

[0057] Genomic DNA of the invention comprises the protein-coding region for a polypeptide of the invention and is also intended to include allelic variants thereof. It is widely understood that, for many genes, genomic DNA is transcribed into RNA transcripts that undergo one or more splicing events wherein intron (i.e., non-coding regions) of the transcripts are removed, or "spliced out." RNA transcripts that can be spliced by alternative mechanisms, and therefore be subject to removal of different RNA sequences but still encode a T1R3 polypeptide, are referred to in the art as splice variants which are embraced by the invention. Splice variants comprehended by the invention therefore are encoded by the same original genomic DNA sequences but arise from distinct mRNA transcripts. Allelic variants are modified forms of a wild-type gene sequence, the modification resulting from recombination during chromosomal

segregation or exposure to conditions which give rise to genetic mutation. Allelic variants, like wild type genes, are naturally occurring sequences (as opposed to non-naturally occurring variants that arise from *in vitro* manipulation).

[0058] The invention also comprehends cDNA that is obtained through reverse transcription of an RNA polynucleotide encoding a T1R3 receptor (conventionally followed by second strand synthesis of a complementary strand to provide a double-stranded DNA).

[0059] One embodiment of the DNA of the invention comprises a double-stranded molecule along with the complementary molecule (the "non-coding strand" or "complement") having a sequence unambiguously deducible from the coding strand according to Watson-Crick base-pairing rules for DNA.

[0060] The present invention includes fragments of nucleotide sequences encoding a T1R3 receptor comprising at least 10, and preferably at least 12, 14, 16, 18, 20, 25, 30, 35, 40, 45, 50, 75, or 100 consecutive nucleotides of a polynucleotide encoding T1R3. polynucleotides of the invention may comprise sequences unique to the T1R3-encoding polynucleotide sequence, and therefore hybridize under highly stringent or moderately stringent conditions only (i.e., "specifically") to polynucleotides encoding T1R3 (or fragments thereof). Polynucleotide fragments of genomic sequences of the invention comprise not only sequences unique to the coding region, but also include fragments of the full-length sequence derived from introns, regulatory regions, and/or other non-translated sequences. Sequences unique to polynucleotides of the invention are recognizable through sequence comparison to other known polynucleotides, and can be identified through use of alignment programs routinely utilized in the art, e.g., those made available in public sequence databases. Such sequences also are recognizable from Southern hybridization analyses to determine the number of fragments of genomic DNA to which a polynucleotide will hybridize. Polynucleotides of the invention can be labeled in a manner that permits their detection, including radioactive, fluorescent, and enzymatic labeling.

[0061] Fragment polynucleotides are particularly useful as probes for detection of full-length or fragments of T1R3 polynucleotides. One or more polynucleotides can be included in kits that are used to detect the presence of a polynucleotide encoding T1R3, or used to detect variations in a polynucleotide sequence encoding T1R3.

[0062] The invention also embraces DNAs encoding T1R3 polypeptides that hybridize under moderately stringent or high stringency conditions to the non-coding strand, or complement, of the polynucleotides.

[0063] Exemplary highly stringent hybridization conditions are as follows: hybridization at 42°C in a hybridization solution comprising 50% formamide, 1% SDS, 1 M NaCl, 10% Dextran sulfate, and washing twice for 30 minutes at 60°C in a wash solution comprising 0.1 X SSC and 1% SDS. It is understood in the art that conditions of equivalent stringency can be achieved through variation of temperature and buffer, or salt concentration as described, for example, in Ausubel et al. (Eds.), PROTOCOLS IN MOLECULAR BIOLOGY, John Wiley & Sons (1994), pp. 6.0.3 to 6.4.10. Modifications in hybridization conditions can be empirically determined or precisely calculated based on the length and the percentage of guanosine/cytosine (GC) base pairing of the probe. The hybridization conditions can be calculated as described, for example, in Sambrook et al., (Eds.), Molecular Cloning: A Laboratory Manual, Cold Spring Harbor Laboratory Press: Cold Spring Harbor, New York (1989), pp. 9.47 to 9.51.

[0064] With the knowledge of the nucleotide sequence information disclosed in the present invention, one skilled in the art can identify and obtain nucleotide sequences which encode T1R3 from different sources (i.e., different tissues or different organisms) through a variety of means well known to the skilled artisan and as disclosed by, for example, Sambrook et al., MOLECULAR CLONING: A LABORATORY MANUAL, Second Edition, Cold Spring Harbor Press, Cold Spring Harbor, NY (1989).

[0065] For example, DNA that encodes T1R3 may be obtained by screening mRNA, cDNA, or genomic DNA with oligonucleotide probes generated from the T1R3 gene sequence information provided herein. Probes may be labeled with a detectable group, such as a fluorescent group, a radioactive atom or a chemiluminescent group in accordance with procedures known to the skilled artisan and used in conventional hybridization assays, as described by, for example, Sambrook *et al*.

[0066] A nucleic acid molecule comprising a T1R3 nucleotide sequence can alternatively be synthesized by use of the polymerase chain reaction (PCR) procedure, with the PCR oligonucleotide primers produced from the nucleotide sequences provided herein. See U.S. Patent Numbers 4,683,195 to Mullis *et al.* and 4,683,202 to Mullis. The PCR reaction provides a method for selectively increasing the concentration of a particular nucleic acid sequence even when that sequence has not been previously purified and is present only in a single copy in a

particular sample. The method can be used to amplify either single- or double-stranded DNA. The essence of the method involves the use of two oligonucleotide probes to serve as primers for the template-dependent, polymerase mediated replication of a desired nucleic acid molecule.

[0067] A wide variety of alternative cloning and *in vitro* amplification methodologies are well known to those skilled in the art. Examples of these techniques are found in, for example, Berger *et al.*, *Guide to Molecular Cloning Techniques*, METHODS IN ENZYMOLOGY 152, Academic Press, Inc., San Diego, CA (Berger), which is incorporated herein by reference in its entirety.

[0068] The polynucleotides of the invention may be used in hybridization techniques known to those skilled in the art, including but not limited to, Northern and Southern blotting and overgo hybridization (see infra). For example, polynucleotide probes of the invention may be used in tissue distribution studies and diagnostic assays. The T1R3 receptor of the invention is likely to be present and active in tissues other than those involved in taste perception. It is therefore likely that the feline T1R3 receptor serves multiple functions in vivo, such as, for example, regulation of amino acid metabolism in addition to taste perception.

[0069] Automated sequencing methods can be used to obtain or verify the nucleotide sequence of T1R3. The nucleotide sequences of the present invention are believed to be accurate. However, as is known in the art, nucleotide sequence obtained by automated methods may contain some errors. Nucleotide sequences determined by automation are typically at least about 90%, more typically at least about 95% to at least about 99.9% identical to the actual nucleotide sequence of a given nucleic acid molecule. The actual sequence may be more precisely determined using manual sequencing methods, which are well known in the art. An error in a sequence which results in an insertion or deletion of one or more nucleotides may result in a frame shift in translation such that the predicted amino acid sequence will differ from that which would be predicted from the actual nucleotide sequence of the nucleic acid molecule, starting at the point of the mutation.

[0070] The nucleic acid molecules of the present invention, and fragments derived therefrom, are useful for screening for restriction fragment length polymorphism (RFLP) associated with certain disorders, as well as for genetic mapping.

[0071] The polynucleotide sequence information provided by the invention makes possible large-scale expression of the encoded polypeptide by techniques well known and routinely practiced in the art.

#### Vectors

[0072] Another aspect of the present invention is directed to vectors, or recombinant expression vectors, comprising any of the nucleic acid molecules described above. Vectors are used herein either to amplify DNA or RNA encoding T1R3 receptor and/or to express DNA which encodes T1R3 receptor. Examples of vectors include, but are not limited to, plasmids, phages, cosmids, episomes, viral particles or viruses, and integratable DNA fragments (i.e., fragments integratable into the host genome by homologous recombination). Examples of viral particles include, but are not limited to, adenoviruses, baculoviruses, parvoviruses, herpesviruses, poxyiruses, adeno-associated viruses, Semliki Forest viruses, vaccinia viruses, and retroviruses. Examples of expression vectors include, but are not limited to, pcDNA3 (Invitrogen) and pSVL (Pharmacia Biotech). Other expression vectors include, but are not limited to, pSPORTTM vectors, pGEM<sup>™</sup> vectors (Promega), pPROEXvectors<sup>™</sup> (LTI, Bethesda, MD), Bluescript<sup>™</sup> (Stratagene),  $pQE^{TM}$ vectors (Qiagen), pSE420TM (Invitrogen), and vectors pYES2<sup>TM</sup>(Invitrogen).

[0073] Expression constructs may comprise T1R3-encoding polynucleotides operably linked to an endogenous or exogenous expression control DNA sequence and a transcription terminator. Expression control DNA sequences include promoters, enhancers, operators, and regulatory element binding sites generally, and are typically selected based on the expression systems in which the expression construct is to be utilized. Promoter and enhancer sequences are generally selected for the ability to increase gene expression, while operator sequences are generally selected for the ability to regulate gene expression. Expression constructs of the invention may also include sequences encoding one or more selectable markers that permit identification of host cells bearing the construct. Expression constructs may also include sequences that facilitate, or promote, homologous recombination in a host cell. Constructs of the invention also may include sequences necessary for replication in a host cell.

[0074] Expression constructs may be utilized for production of an encoded protein, but may also be utilized simply to amplify a T1R3-encoding polynucleotide sequence. In some embodiments, the vector is an expression vector wherein a polynucleotide of the invention is operably linked to a polynucleotide comprising an expression control sequence. Autonomously

replicating recombinant expression constructs such as plasmid and viral DNA vectors incorporating polynucleotides of the invention are also provided. Some expression vectors are replicable DNA constructs in which a DNA sequence encoding a T1R3 receptor is operably linked or connected to suitable control sequence(s) capable of effecting the expression of the receptor in a suitable host. Amplification vectors do not require expression control domains, but rather need only the ability to replicate in a host, such as conferred by an origin of replication, and a selection gene to facilitate recognition of transformants. The need for control sequences in the expression vector will vary depending upon the host selected and the transformation method chosen. Control sequences include a transcriptional promoter, an optional operator sequence to control transcription, a sequence encoding suitable mRNA ribosomal binding, and sequences which control the termination of transcription and translation.

[0075] Vectors of the invention may contain a promoter that is recognized by the host organism. The promoter sequences of the present invention may be prokaryotic, eukaryotic, or viral. Examples of suitable prokaryotic sequences include the P<sub>R</sub> and P<sub>L</sub> promoters of bacteriophage lambda (The Bacteriophage Lambda, Hershey, A. D., Ed., Cold Spring Harbor Press, Cold Spring Harbor, NY (1973), which is incorporated herein by reference in its entirety; Lambda II, Hendrix, R. W., Ed., Cold Spring Harbor Press, Cold Spring Harbor, NY (1980), which is incorporated herein by reference in its entirety), the trp, recA, heat shock, and lacZ promoters of *E. coli*, and the SV40 early promoter (Benoist *et al. Nature*, 1981, 290, 304-310), which is incorporated herein by reference in its entirety. Additional promoters include, but are not limited to, mouse mammary tumor virus, long terminal repeat of human immunodeficiency virus, maloney virus, cytomegalovirus immediate early promoter, Epstein Barr virus, Rous sarcoma virus, human actin, human myosin, human hemoglobin, human muscle creatine, and human metallothionein.

[0076] Additional regulatory sequences can also be included in vectors of the invention. Examples of suitable regulatory sequences are represented by the Shine-Dalgarno of the replicase gene of the phage MS-2 and of the gene cII of bacteriophage lambda. The Shine-Dalgarno sequence may be directly followed by DNA encoding a T1R3 receptor, resulting in the expression of the mature protein.

[0077] Moreover, suitable expression vectors can include an appropriate marker that allows the screening of transformed host cells. The transformation of the selected host is carried out using any one of the various techniques well known to the expert in the art and described in Sambrook et al., supra.

[0078] An origin of replication or autonomously replicating sequence (ARS) can also be provided either by construction of the vector to include an exogenous origin or may be provided by the host cell chromosomal replication mechanism. If the vector is integrated into the host cell chromosome, the latter may be sufficient. Alternatively, rather than using vectors which contain viral origins of replication, one skilled in the art can transform mammalian cells by the method of co-transformation with a selectable marker and T1R3 DNA. An example of a suitable marker is dihydrofolate reductase (DHFR) or thymidine kinase (see, U.S. Patent No. 4,399,216).

[0079] Additional regulatory sequences that may be included in the polynucleotides of the invention include secretion signals which allow the encoded polypeptide to cross and/or lodge in cell membranes, or be secreted from the cell.

Nucleotide sequences encoding T1R3 may be recombined with vector DNA in accordance with conventional techniques, including blunt-ended or staggered-ended termini for ligation, restriction enzyme digestion to provide appropriate termini, filling in of cohesive ends as appropriate, alkaline phosphatase treatment to avoid undesirable joining, and ligation with appropriate ligases. Techniques for such manipulation are disclosed by Sambrook *et al.*, *supra* and are well known in the art. Methods for construction of mammalian expression vectors are disclosed in, for example, Okayama *et al.*, *Mol. Cell. Biol.*, 1983, 3, 280, Cosman *et al.*, *Mol. Immunol.*, 1986, 23, 935, Cosman *et al.*, *Nature*, 1984, 312, 768, EP-A-0367566, and WO 91/18982, each of which is incorporated herein by reference in its entirety.

#### Host cells

[0081] According to another aspect of the invention, host cells are provided, including prokaryotic and eukaryotic cells, comprising a polynucleotide of the invention (or vector of the invention) in a manner that permits expression of the encoded T1R3 polypeptide. Polynucleotides of the invention may be introduced into the host cell as part of a circular plasmid, or as linear DNA comprising an isolated protein-coding region or a viral vector. Methods for introducing DNA into the host cell that are well known and routinely practiced in the art include transformation, transfection, electroporation, nuclear injection, or fusion with carriers such as liposomes, micelles, ghost cells, and protoplasts. Expression systems of the invention include bacterial, yeast, fungal, plant, insect, invertebrate, vertebrate, and mammalian cell systems.

[0082] The invention provides host cells that are transformed or transfected (stably or transiently) with polynucleotides of the invention or vectors of the invention. As stated above, such host cells are useful for amplifying the polynucleotides and also for expressing a T1R3 polypeptide or fragment thereof encoded by the polynucleotide.

[0083] In still another related embodiment, the invention provides a method for producing a T1R3 polypeptide (or fragment thereof) comprising the steps of growing a host cell of the invention in a nutrient medium and isolating the polypeptide or variant thereof from the cell or the medium. Because the T1R3 receptor is a membrane-spanning polypeptide, it will be appreciated that, for some applications, such as certain activity assays, the preferable isolation may involve isolation of cell membranes containing the polypeptide embedded therein, whereas for other applications a more complete isolation may be preferable.

[0084] According to some aspects of the present invention, transformed host cells having an expression vector comprising any of the nucleic acid molecules described above are provided. Expression of the nucleotide sequence occurs when the expression vector is introduced into an appropriate host cell. Suitable host cells for expression of the polypeptides of the invention include, but are not limited to, prokaryotes, yeast, and eukaryotes. If a prokaryotic expression vector is employed, then the appropriate host cell would be any prokaryotic cell capable of expressing the cloned sequences. Suitable prokaryotic cells include, but are not limited to, bacteria of the genera Escherichia, Bacillus, Salmonella, Pseudomonas, Streptomyces, and Staphylococcus.

[0085] If a eukaryotic expression vector is employed, then the appropriate host cell would be any eukaryotic cell capable of expressing the cloned sequence. Eukaryotic cells may be cells of higher eukaryotes. Suitable eukaryotic cells include, but are not limited to, non-human mammalian tissue culture cells and human tissue culture cells. Host cells include, but are not limited to, insect cells, HeLa cells, Chinese hamster ovary cells (CHO cells), African green monkey kidney cells (COS cells), human HEK-293 cells, and murine 3T3 fibroblasts. Propagation of such cells in cell culture has become a routine procedure (see, TISSUE CULTURE, Academic Press, Kruse and Patterson, eds. (1973), which is incorporated herein by reference in its entirety).

[0086] In addition, a yeast host may be employed as a host cell. Yeast cells include, but are not limited to, the genera Saccharomyces, Pichia, and Kluveromyces. Yeast hosts may be S. cerevisiae and P. pastoris. Yeast vectors may contain an origin of replication sequence from a

2T yeast plasmid, an autonomously replication sequence (ARS), a promoter region, sequences for polyadenylation, sequences for transcription termination, and a selectable marker gene. Shuttle vectors for replication in both yeast and *E. coli* are also included herein.

[0087] Alternatively, insect cells may be used as host cells. In some embodiments, the polypeptides of the invention are expressed using a baculovirus expression system (see, Luckow et al., Bio/Technology, 1988, 6, 47; Baculovirus Expression Vectors: A Laboratory Manual, O'Reilly et al. (Eds.), W.H. Freeman and Company, New York, 1992; and U.S. Patent No. 4,879,236, each of which is incorporated herein by reference in its entirety). In addition, the MAXBAC<sup>TM</sup> complete baculovirus expression system (Invitrogen) can, for example, be used for production in insect cells.

Host cells of the invention are a valuable source of immunogen for development of [8800] antibodies specifically immunoreactive with the T1R3 receptor. Host cells of the invention also are useful in methods for the large-scale production of T1R3 polypeptides wherein the cells are grown in a suitable culture medium and the desired polypeptide products are isolated from the cells, or from the medium in which the cells are grown, by purification methods known in the art, e.g., conventional chromatographic methods including immunoaffinity chromatography, receptor hydrophobic interaction chromatography. lectin affinity affinity chromatography, chromatography, size exclusion filtration, cation or anion exchange chromatography, high pressure liquid chromatography (HPLC), reverse phase HPLC, and the like. Still other methods of purification include those methods wherein the desired protein is expressed and purified as a fusion protein having a specific tag, label, or chelating moiety that is recognized by a specific binding partner or agent. The purified protein can be cleaved to yield the desired protein, or can be left as an intact fusion protein. Cleavage of the fusion component may produce a form of the desired protein having additional amino acid residues as a result of the cleavage process.

[0089] Knowledge of the feline T1R3 nucleotide sequence allows for modification of cells to permit, or increase, expression of endogenous receptor. Cells can be modified (e.g., by homologous recombination) to provide increased expression by replacing, in whole or in part, the naturally occurring T1R3 promoter with all or part of a heterologous promoter so that the cells express the receptor at higher or lower levels. The heterologous promoter is inserted in such a manner that it is operably linked to endogenous T1R3 coding sequence. (See, for example, PCT International Publication No. WO 94/12650, PCT International Publication No. WO 92/20808, and PCT International Publication No. WO 91/09955.) It is also contemplated that, in addition to heterologous promoter DNA, amplifiable marker DNA (e.g., ada, dhfr, and

the multifunctional CAD gene which encodes carbamoyl phosphate synthase, aspartate transcarbamylase, and dihydroorotase) and/or intron DNA may be inserted along with the heterologous promoter DNA. If linked to the T1R3 coding sequence, amplification of the marker DNA by standard selection methods results in co-amplification of the T1R3 coding sequences in the cells.

#### Knock-out and transplacement animals

[0090] The DNA sequence information provided by the present invention also makes possible the development (e.g., by homologous recombination strategies; see Capecchi, Science 244:1288-1292 (1989), which is incorporated herein by reference) of transgenic animals, including, for example, animals that fail to express functional T1R3 ("knock-out") or that express a variant thereof ("transplacement"). Such animals (especially small laboratory animals such as rats, rabbits, mice, and cats) are useful as models for studying the in vivo activities of T1R3 and modulators of T1R3.

#### Antisense and siRNA

[0091] Also encompassed by the invention are antisense and short interfering polynucleotides that recognize and hybridize to polynucleotides encoding T1R3. Full-length and fragment antisense polynucleotides are provided. Fragment antisense molecules of the invention include (i) those that specifically recognize and hybridize to T1R3 RNA (as determined by sequence comparison of DNA encoding T1R3 to DNA encoding other known molecules). Identification of sequences unique to T1R3 encoding polynucleotides can be deduced through use of any publicly available sequence database, and/or through use of commercially available sequence comparison programs. After identification of the desired sequences, isolation through restriction digestion or amplification using any of the various polymerase chain reaction techniques well known in the art can be performed. Antisense polynucleotides are particularly relevant to regulation of expression of T1R3 receptor by those cells expressing T1R3 mRNA.

[0092] Antisense nucleic acids (preferably 10 to 30 base-pair oligonucleotides) capable of specifically binding to T1R3 expression control sequences or T1R3 RNA are introduced into cells (e.g., by a viral vector or colloidal dispersion system such as a liposome). The antisense nucleic acid binds to the target nucleotide sequence in the cell and prevents transcription and/or translation of the target sequence. Phosphorothioate and methylphosphonate antisense oligonucleotides are specifically contemplated for therapeutic use by the invention. Locked

nucleic acids are also specifically contemplated for therapeutic use by the present invention. (See, for example, Wahlestedt et al., Proc. Natl. Acad. Sci. USA, 97(10), 5633-5638 (2000), which is incorporated by reference in its entirety) The antisense oligonucleotides may be further modified by adding poly-L-lysine, transferrin polylysine, or cholesterol moieties at their 5' end. Suppression of T1R3 expression at either the transcriptional or translational level is useful to generate cellular or animal models for diseases/conditions characterized by aberrant T1R3 expression.

[0093] Antisense oligonucleotides, or fragments of nucleotide sequence of SEQ ID NO:1, or sequences complementary or homologous thereto, derived from the nucleotide sequences of the present invention encoding T1R3 receptor are useful as diagnostic tools for probing gene expression in various tissues. For example, tissue can be probed *in situ* with oligonucleotide probes carrying detectable groups by conventional autoradiography techniques to investigate native expression of this enzyme or pathological conditions relating thereto. Antisense oligonucleotides may be directed to regulatory regions of a T1R3 nucleotide sequence, or mRNA corresponding thereto, including, but not limited to, the initiation codon, TATA box, enhancer sequences, and the like.

[0094] Those of skill in the art recognize that the antisense oligonucleotides that inhibit the expression and/or biological activity of a T1R3 receptor may be predicted using any gene encoding a T1R3 receptor. Specifically, antisense nucleic acid molecules comprise a sequence preferably complementary to at least about 5, 10, 15, 20, 25, 30, 35, 40, 45, 50, 100, 250 or 500 nucleotides or an entire T1R3 receptor gene sequence. The antisense oligonucleotides may comprise a sequence complementary to about 15 consecutive nucleotides of the coding strand of the T1R3 receptor-encoding sequence.

[0095] In one embodiment, an antisense nucleic acid molecule is antisense to a "coding region" of the coding strand of a nucleotide sequence encoding a T1R3 protein. The coding strand may also include regulatory regions of the T1R3 sequence. The term "coding region" refers to the region of the nucleotide sequence comprising codons which are translated into amino acid residues. In another embodiment, the antisense nucleic acid molecule is antisense to a "noncoding region" of the coding strand of a nucleotide sequence encoding a T1R3 protein. The term "noncoding region" refers to 5' and 3' sequences which flank the coding region that are not translated into amino acids (i.e., also referred to as 5' and 3' untranslated regions (UTR)).

[0096] Antisense oligonucleotides may be directed to regulatory regions of a nucleotide sequence encoding a T1R3 protein, or mRNA corresponding thereto, including, but not limited to, the initiation codon, TATA box, enhancer sequences, and the like. Given the coding strand sequences provided herein, antisense nucleic acids of the invention can be designed according to the rules of Watson and Crick or Hoogsteen base pairing. The antisense nucleic acid molecule can be complementary to the entire coding region of a T1R3 mRNA, but also may be an oligonucleotide that is antisense to only a portion of the coding or noncoding region of the mRNA. For example, the antisense oligonucleotide can be complementary to the region surrounding the translation start site of an mRNA. An antisense oligonucleotide can be, for example, about 5, 10, 15, 20, 25, 30, 35, 40, 45 or 50 nucleotides in length.

[0097] Another means to inhibit the activity of a T1R3 receptor according to the invention is via RNA interference (RNAi) (see e.g., Elbashir et al., Nature, 411:494-498 (2001); Elbashir et al., Genes Development, 15:188-200 (2001)). RNAi is the process of sequence-specific, post-transcriptional gene silencing, initiated by double-stranded RNA (dsRNA) that is homologous in sequence to the silenced gene (e.g., is homologous in sequence to the sequence of a T1R3, for example but not limited to the sequence as set forth in SEQ ID NO:1). siRNA-mediated silencing is thought to occur post-transcriptionally and/or transcriptionally. For example, siRNA duplexes may mediate post-transcriptional gene silencing by reconstitution of siRNA-protein complexes (siRNPs), which guide mRNA recognition and targeted cleavage.

[0098] Accordingly, another form of a T1R3 inhibitory compound of the invention is a short interfering RNA (siRNA) directed against a T1R3-encoding sequence. Exemplary siRNAs are siRNA duplexes (for example, 10-25, preferably 20, 21, 22, 23, 24, or 25 residues in length) having a sequence homologous or identical to a fragment of the T1R3 sequence set forth as SEQ ID NO:1 and having a symmetric 2-nucleotide 3'-overhang. The 2-nucleotide 3' overhang may be composed of (2'-deoxy) thymidine because it reduces costs of RNA synthesis and may enhance nuclease resistance of siRNAs in the cell culture medium and within transfected cells. Substitution of uridine by thymidine in the 3' overhang is also well tolerated in mammalian cells, and the sequence of the overhang appears not to contribute to target recognition.

#### **Polypeptides**

[0099] The invention also provides purified and isolated mammalian T1R3 receptor polypeptides encoded by a polynucleotide of the invention. Some embodiments include a feline T1R3 polypeptide comprising the amino acid sequence of SEQ ID NO:2, or fragments thereof

comprising an epitope specific to the polypeptide. A reference to "epitope specific to" or "polypeptide-specific epitope," or variations thereof, indicates that a portion of the T1R3 receptor or amino acid sequence is recognizable by an antibody that is specific for the T1R3 or amino acid sequence.

[0100] Included within the scope of the invention are polypeptides encoded by feline allelic variants of T1R3. The allelic variants of the T1R3 receptor of the invention may modify the taste perception of a mammal, such as a cat, to a taste stimulus. Such functional amino acid sequence modifications may account for differences in intraspecies (e.g., breed-specific) taste perception.

[0101] Extracellular epitopes are useful for generating and screening for antibodies and other binding compounds that bind to a T1R3 receptor. Thus, in another embodiment, the invention provides a purified and isolated polypeptide comprising at least one extracellular domain of the T1R3 receptor. Also included is a polypeptide comprising a T1R3 receptor fragment selected from the group consisting of an extracellular domain of T1R3 (residues 1-571, 628-641, 705-730, and 787-794 of SEQ ID NO:2), a transmembrane domain of T1R3 (residues 572-594, 610-627, 642-664, 681-704, 731-754, 767-780, and 795-812 of SEQ ID NO:2), and an intracellular domain of T1R3 (residues 595-609, 665-680, 755-766, and 813-865 of SEQ ID NO:2). Polypeptide fragments of the invention may be continuous portions of the native receptor. However, it will also be appreciated that knowledge of the T1R3 gene and protein sequences as provided herein permits recombination of various domains that are not contiguous in the native protein.

[0102] The invention embraces polypeptides that preferably have at least 99%, at least 95%, at least 90%, at least 85%, at least 75%, at least 74%, at least 73%, at least 72%, at least 71%, at least 70%, at least 65%, at least 60%, at least 55% or at least 50% identity and/or homology to the polypeptide of the invention.

[0103] Polypeptides of the invention may be isolated from natural cell sources or may be chemically synthesized, but are preferably produced by recombinant procedures involving host cells of the invention. Use of mammalian host cells is expected to provide for such post-translational modifications (e.g., glycosylation, truncation, lipidation, and phosphorylation) as may be needed to confer optimal biological activity on recombinant expression products of the invention.

[0104] The invention also embraces variant T1R3 polypeptides. In one example, insertion variants are provided wherein one or more amino acid residues supplement a T1R3 amino acid sequence such as SEQ ID NO:2. Insertions may be located at either or both termini of the protein, or may be positioned within internal regions of the amino acid sequence. Insertional variants with additional residues at either or both termini can include, for example, fusion proteins and proteins including amino acid tags or labels.

[0105] Insertion variants include T1R3 polypeptides wherein one or more amino acid residues are added to a biologically active fragment thereof.

[0106] The invention also embraces T1R3 variants having additional amino acid residues that result from use of specific expression systems. For example, use of commercially available vectors that express a desired polypeptide as part of a glutathione-S-transferase (GST) fusion product provides the desired polypeptide having an additional glycine residue at position -1 after cleavage of the GST component from the desired polypeptide. Variants that result from expression in other vector systems are also contemplated.

[0107] In another aspect, the invention provides deletion variants wherein one or more amino acid residues in a T1R3 polypeptide are removed. Deletions can be effected at one or both termini of the T1R3 polypeptide, or with removal of one or more non-terminal amino acid residues of T1R3. Deletion variants, therefore, include all fragments of a T1R3 polypeptide.

[0108] The invention also embraces polypeptide fragments that maintain biological (e.g., ligand binding) and/or immunological properties of a T1R3 polypeptide.

[0109] As used in the present invention, polypeptide fragments preferably comprise at least 10, 15, 20, 25, 30, 35, 40, 45, or 50 consecutive amino acids of SEQ ID NO:2. Some polypeptide fragments display antigenic properties unique to, or specific for, feline T1R3. Fragments of the invention having the desired biological and immunological properties can be prepared by any of the methods well known and routinely practiced in the art.

[0110] In still another aspect, the invention provides substitution variants of T1R3 polypeptides. Substitution variants include those polypeptides wherein one or more amino acid residues of a T1R3 polypeptide are removed and replaced with alternative residues. In one aspect, the substitutions are conservative in nature; however, the invention embraces substitutions that are also non-conservative. Conservative substitutions for this purpose may be defined as set out in Tables 1, 2, or 3 below.

[0111] Variant polypeptides include those wherein conservative substitutions have been introduced by modification of polynucleotides encoding polypeptides of the invention. Amino acids can be classified according to physical properties and contribution to secondary and tertiary protein structure. A conservative substitution is recognized in the art as a substitution of one amino acid for another amino acid that has similar properties. Exemplary conservative substitutions are set out in Table 1 (from WO 97/09433, page 10, published March 13, 1997 (PCT/GB96/02197, filed 9/6/96), immediately below.

Table 1
Conservative Substitutions I

SIDE CHAIN	
<b>CHARACTERISTIC</b>	AMINO ACID
Aliphatic	
Non-polar	GAP
	ILV
Polar - uncharged	CSTM
	NQ
Polar - charged	DE
Ü	KR
Aromatic	HFWY
Other	NQDE

Alternatively, conservative amino acids can be grouped as described in Lehninger, [BIOCHEMISTRY, Second Edition; Worth Publishers, Inc. NY, NY (1975), pp.71-77] as set out in Table 2, below.

Table 2
Conservative Substitutions II

SIDE CHAIN CHARACTERISTIC	AMINO ACID
Non-polar (hydrophobic)	
A. Aliphatic:	ALIVP
B. Aromatic:	F W
C. Sulfur-containing:	M
D. Borderline:	G
Uncharged-polar	
A. Hydroxyl:	STY
B. Amides:	ΝQ
C. Sulfhydryl:	C
D. Borderline:	G
Positively Charged (Basic):	KRH
Negatively Charged (Acidic):	DE

As still another alternative, exemplary conservative substitutions are set out in Table 3, below.

Table 3

#### **Conservative Substitutions III**

Original Residue	<b>Exemplary Substitution</b>
Ala (A)	Val, Leu, Ile
Arg (R)	Lys, Gln, Asn
Asn (N)	Gln, His, Lys, Arg
Asp (D)	Glu
Cys (C)	Ser
Gln (Q)	Asn
Glu (E)	Asp
His (H)	Asn, Gln, Lys, Arg
Ile (I)	Leu, Val, Met, Ala, Phe,
Leu (L)	Ile, Val, Met, Ala, Phe
Lys (K)	Arg, Gln, Asn
Met (M)	Leu, Phe, Ile
Phe (F)	Leu, Val, Ile, Ala
Pro (P)	Gly
Ser (S)	Thr
Thr (T)	Ser
Trp (W)	Tyr
Tyr (Y)	Trp, Phe, Thr, Ser
Val (V)	Ile, Leu, Met, Phe, Ala

[0112] It should be understood that the definition of polypeptides of the invention is intended to include polypeptides bearing modifications other than insertion, deletion, or substitution of amino acid residues. By way of example, the modifications may be covalent in nature, and include for example, chemical bonding with polymers, lipids, other organic, and inorganic moieties. Such derivatives may be prepared to increase circulating half-life of a polypeptide, or may be designed to improve the targeting capacity of the polypeptide for desired cells, tissues, or organs. Similarly, the invention further embraces T1R3 polypeptides that have been covalently modified to include one or more water-soluble polymer attachments such as polyethylene glycol, polyoxyethylene glycol, or polypropylene glycol. Variants that display ligand binding properties of native T1R3 and are expressed at higher levels, as well as variants that provide for constitutively active receptors, are particularly useful in assays of the invention; the variants are also useful in providing cellular, tissue and animal models of diseases/conditions characterized by aberrant T1R3 activity.

[0113] In a related embodiment, the present invention provides compositions comprising purified polypeptides of the invention. Some compositions comprise, in addition to the

polypeptide of the invention, a pharmaceutically acceptable (i.e., sterile and non-toxic) liquid, semisolid, or solid diluent that serves as a pharmaceutical vehicle, excipient, or medium. Any diluent known in the art may be used. Exemplary diluents include, but are not limited to, water, saline solutions, polyoxyethylene sorbitan monolaurate, magnesium stearate, methyl- and propylhydroxybenzoate, talc, alginates, starches, lactose, sucrose, dextrose, sorbitol, mannitol, glycerol, calcium phosphate, mineral oil, and cocoa butter.

[0114] Variants that display ligand-binding properties of native T1R3 and are expressed at higher levels, as well as variants that provide for constitutively active receptors, are particularly useful in assays of the invention; the variants are also useful in assays of the invention and in providing cellular, tissue and animal models of diseases/conditions characterized by aberrant T1R3 activity.

#### **Antibodies**

Also included in the present invention are antibodies (e.g., monoclonal and polyclonal antibodies, single chain antibodies, chimeric antibodies, bifunctional/bispecific antibodies, humanized antibodies, human antibodies, felinized antibodies, feline antibodies, and complementary determining region (CDR)-grafted antibodies, including compounds which include CDR sequences which specifically recognize a polypeptide of the invention) specific for T1R3 receptor or fragments thereof. Antibody fragments, including Fab, Fab', F(ab')2, and F<sub>v</sub>, are also provided by the invention. The term "specific for," when used to describe antibodies of the invention, indicates that the variable regions of the antibodies of the invention recognize and bind T1R3 polypeptides, preferably exclusively (i.e., are able to distinguish T1R3 polypeptides from other known polypeptides by virtue of measurable differences in binding affinity, despite the possible existence of localized sequence identity, homology, or similarity between T1R3 and such polypeptides). It will be understood that specific antibodies may also interact with other proteins (for example, S. aureus protein A or other antibodies in ELISA techniques) through interactions with sequences outside the variable region of the antibodies, and, in particular, in the constant region of the molecule. Screening assays to determine binding specificity of an antibody of the invention are well known and routinely practiced in the art. For a comprehensive discussion of such assays, see Harlow et al. (Eds.), ANTIBODIES A LABORATORY MANUAL; Cold Spring Harbor Laboratory; Cold Spring Harbor, NY (1988), Chapter 6. Antibodies that recognize and bind fragments of the T1R3 polypeptides of the invention are also contemplated, provided that the antibodies are specific for T1R3 polypeptides. Antibodies of the invention can be produced using any method well known and routinely practiced in the art.

[0116] The invention provides an antibody that is specific for the feline T1R3 of the invention. Antibodies that can be generated from polypeptides that have previously been described in the literature and that are capable of fortuitously cross-reacting with T1R3 (e.g., due to the fortuitous existence of a similar epitope in both polypeptides) are considered "cross-reactive" antibodies. Such cross-reactive antibodies are not antibodies that are "specific" for T1R3 receptor. The determination of whether an antibody is specific for T1R3 or is cross-reactive with another known receptor is made using any of several assays, such as Western blotting assays, that are well known in the art. For identifying cells that express T1R3 and also for modulating T1R3-ligand binding activity, antibodies that specifically bind to an extracellular epitope of the T1R3 may be used.

[0117] In some variations, the invention provides monoclonal antibodies. Hybridomas that produce such antibodies also are intended as aspects of the invention. In yet another variation, the invention provides a felinized antibody. Felinized antibodies are useful for *in vivo* therapeutic indications.

[0118] In another variation, the invention provides a cell-free composition comprising polyclonal antibodies, wherein at least one of the antibodies is an antibody of the invention specific for T1R3. Antisera isolated from an animal is an exemplary composition, as is a composition comprising an antibody fraction of an antisera that has been resuspended in water or in another diluent, excipient, or carrier.

[0119] In still another related embodiment, the invention provides an anti-idiotypic antibody specific for an antibody that is specific for T1R3.

[0120] It is well known that antibodies contain relatively small antigen binding domains that can be isolated chemically or by recombinant techniques. Such domains are useful T1R3 binding molecules themselves, and also may be reintroduced into other antibodies or fused to toxins or other polypeptides. Thus, in still another embodiment, the invention provides a polypeptide comprising a fragment of a T1R3-specific antibody, wherein the fragment and the polypeptide bind to the T1R3 receptor. By way of non-limiting example, the invention provides polypeptides that are single chain antibodies and CDR-grafted antibodies.

[0121] Non-feline antibodies may be felinized by any of the methods known in the art. In one method, the non-feline CDRs are inserted into a feline antibody or consensus antibody framework sequence. Similarly, non-human antibodies may be humanized by methods known in

the art. In one embodiment, non-human CDRs are inserted into a human antibody or consensus antibody framework sequence. Further changes can then be introduced into the antibody framework to modulate affinity or immunogenicity.

[0122] Antibodies of the invention are useful for, e.g., therapeutic purposes (such as by modulating activity of T1R3), diagnostic purposes (such as detecting or quantitating T1R3 activity), and also for purification of T1R3. Kits comprising an antibody of the invention for any of the purposes described herein are also included within the scope of the invention. In general, a kit of the invention preferably includes a control antigen for which the antibody is immunospecific.

## Compositions

[0123] Mutations in the T1R3 gene that result in loss of normal function of the T1R3 gene product underlie some T1R3-related disease states. The invention comprehends gene and peptide therapy, for example, to restore T1R3 activity to treat those disease states. Delivery of a functional T1R3 gene to appropriate cells is effected ex vivo, in situ, or in vivo by use of vectors, and more particularly viral vectors (e.g., adenovirus, adeno-associated virus, or a retrovirus), or ex vivo by use of physical DNA transfer methods (e.g., liposomes or chemical treatments). See, for example, Anderson, Nature, supplement to vol. 392, No. 6679, pp.25-20 (1998). For additional reviews of gene therapy technology see Friedmann, Science, 244: 1275-1281 (1989); Verma, Scientific American: 68-84 (1990); and Miller, Nature, 357: 455-460 (1992). Alternatively, it is contemplated that in other disease states, preventing the expression of, or inhibiting the activity of, T1R3 will be useful in treatment. It is contemplated that antisense therapy or gene therapy could be applied to negatively regulate the expression of T1R3.

[0124] Another aspect of the present invention is directed to compositions, including pharmaceutical compositions, comprising any of the nucleic acid molecules or recombinant expression vectors described above and an acceptable carrier or diluent. The carrier or diluent may be pharmaceutically acceptable. Suitable carriers are described in the most recent edition of *Remington's Pharmaceutical Sciences*, A. Osol, a standard reference text in this field, which is incorporated herein by reference in its entirety. Examples of such carriers or diluents include, but are not limited to, water, saline, Ringer's solution, dextrose solution, and 5% serum albumin. Liposomes and nonaqueous vehicles such as fixed oils may also be used. The formulations may be sterilized by commonly used techniques.

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[0125] Also within the scope of the invention are compositions comprising polypeptides, polynucleotides, or antibodies of the invention that have been formulated with, e.g., a pharmaceutically acceptable carrier.

[0126] The invention also provides methods of using antibodies of the invention. For example, the invention provides a method for modulating ligand-binding of a T1R3 receptor comprising the step of contacting the receptor with an antibody specific for the T1R3 polypeptide, under conditions wherein the antibody binds the receptor.

#### Methods of identifying ligands and modulators

[0127] The invention also provides assays to identify compounds that bind and/or modulate T1R3. A "T1R3 binding partner" is a compound that directly or indirectly binds a T1R3 polypeptide of the invention. One assay of the invention comprises the steps of: (a) contacting T1R3 with a compound suspected of binding T1R3; and (b) measuring binding between the compound and T1R3. In one variation, the composition comprises a cell expressing T1R3 on its surface. In another variation, isolated T1R3 or cell membranes comprising T1R3 are employed. The binding may be measured directly, e.g., by using a labeled compound, or may be measured indirectly. Compounds identified as binding T1R3 receptor may be further tested in other assays including, but not limited to, T1R3 activity assays and/or in vivo models, in order to confirm or quantitate their activity.

[0128] Specific binding molecules, including natural ligands and synthetic compounds, can be identified or developed using isolated or recombinant T1R3 products, T1R3 variants, or preferably, cells expressing such products. Binding partners are useful for purifying T1R3 products and detection or quantification of T1R3 products in fluid and tissue samples using known immunological procedures. Binding molecules are also manifestly useful in modulating (i.e., blocking, inhibiting or stimulating) biological activities of T1R3, especially those activities involved in signal transduction.

[0129] The DNA and amino acid sequence information provided by the present invention also makes possible identification of binding partner compounds with which a T1R3 polypeptide or polynucleotide will interact. Methods to identify binding partner compounds include solution assays, *in vitro* assays wherein T1R3 polypeptides are immobilized, and cell-based assays. Identification of binding partner compounds of T1R3 polypeptides provides candidates for

therapeutic or prophylactic intervention in pathologies associated with T1R3 normal and aberrant biological activity.

[0130] The invention includes several assay systems for identifying T1R3-binding partners. In solution assays, methods of the invention comprise the steps of (a) contacting a T1R3 polypeptide with one or more candidate binding partner compounds and (b) identifying the compounds that bind to the T1R3 polypeptide. Identification of the compounds that bind the T1R3 polypeptide can be achieved by isolating the T1R3 polypeptide/binding partner complex, and separating the binding partner compound from the T1R3 polypeptide. An additional step of characterizing the physical, biological, and/or biochemical properties of the binding partner compound is also comprehended in another embodiment of the invention. In one aspect, the T1R3 polypeptide/binding partner complex is isolated using an antibody immunospecific for either the T1R3 polypeptide or the candidate binding partner compound.

[0131] In still other embodiments, either the T1R3 polypeptide or the candidate binding partner compound comprises a label or tag that facilitates its isolation, and methods of the invention to identify binding partner compounds include a step of isolating the T1R3 polypeptide/binding partner complex through interaction with the label or tag. An exemplary tag of this type is a poly-histidine sequence, generally around six histidine residues, that permits isolation of a compound so labeled using nickel chelation. Other labels and tags, such as the FLAG® tag (Eastman Kodak, Rochester, NY), well known and routinely used in the art, are embraced by the invention.

[0132] In one variation of an *in vitro* assay, the invention provides a method comprising the steps of (a) contacting an immobilized T1R3 polypeptide with a candidate binding partner compound and (b) detecting binding of the candidate compound to the T1R3 polypeptide. In an alternative embodiment, the candidate binding partner compound is immobilized and binding of T1R3 is detected. Immobilization is accomplished using any of the methods well known in the art, including covalent bonding to a support, a bead, or a chromatographic resin, as well as non-covalent, high affinity interactions such as antibody binding, or use of streptavidin/biotin binding wherein the immobilized compound includes a biotin moiety. The support may, for example, be formulated into a feline-specific electronic tongue. Detection of binding can be accomplished (i) using a radioactive label on the compound that is not immobilized, (ii) using a fluorescent label on the non-immobilized compound, (iv) using a label on the non-immobilized compound that excites a

fluorescent support to which the immobilized compound is attached, as well as other techniques well known and routinely practiced in the art.

[0133] The invention also provides cell-based assays to identify binding partner compounds of a T1R3 polypeptide. In one embodiment, the invention provides a method comprising the steps of contacting a T1R3 polypeptide expressed on the surface of a cell with a candidate binding partner compound and detecting binding of the candidate binding partner compound to the T1R3 polypeptide. In some embodiments, the detection comprises detecting physiological event in the cell caused by the binding of the molecule.

Another aspect of the present invention is directed to methods of identifying [0134] compounds that bind to either T1R3 or nucleic acid molecules encoding T1R3, comprising contacting T1R3, or a nucleic acid molecule encoding the same, with a compound, and determining whether the compound binds T1R3 or a nucleic acid molecule encoding the same. Binding can be determined by binding assays which are well known to the skilled artisan, including, but not limited to, gel-shift assays, Western blots, radiolabeled competition assay, phage-based expression cloning, co-fractionation by chromatography, co-precipitation, crosslinking, interaction trap/two-hybrid analysis, southwestern analysis, ELISA, and the like, which are described in, for example, CURRENT PROTOCOLS IN MOLECULAR BIOLOGY, 1999, John Wiley & Sons, NY, which is incorporated herein by reference in its entirety. The compounds to be screened include (which may include compounds which are suspected to bind T1R3, or a nucleic acid molecule encoding the same), but are not limited to, extracellular, intracellular, biological, or chemical origin. The methods of the invention also embrace ligands, especially neuropeptides, that are attached to a label, such as a radiolabel (e.g., 125I, 35S, 32P, 33P, 3H), a fluorescence label, a chemiluminescent label, an enzymic label, and an immunogenic label. Modulators falling within the scope of the invention include, but are not limited to, non-peptide molecules such as non-peptide mimetics, non-peptide allosteric effectors, and peptides. The TIR3 polypeptide or polynucleotide employed in such a test may either be free in solution, attached to a solid support, borne on a cell surface or located intracellularly, or associated with a portion of a cell. One skilled in the art can, for example, measure the formation of complexes between T1R3 and the compound being tested. Alternatively, one skilled in the art can examine the diminution in complex formation between T1R3 and its substrate caused by the compound being tested. In some embodiments of the invention, the recognition sites of the T1R3 receptor are coupled with a monitoring system, either electrical or optical. An appropriate chemical stimulus can bind to the receptor's ligand binding domain, changing the receptor conformation to a degree that the coupled electronics or optical changes can be observed on a read-out. Such a device could be developed into a feline-specific electronic tongue, for example.

[0135] In another embodiment of the invention, high throughput screening for compounds having suitable binding affinity to T1R3 receptor is employed. Briefly, large numbers of different small peptide test compounds are synthesized on a solid substrate. The peptide test compounds are contacted with T1R3 and washed. Bound T1R3 is then detected by methods well known in the art. Purified polypeptides of the invention can also be coated directly onto plates for use in the aforementioned drug screening techniques. In addition, non-neutralizing antibodies can be used to capture the protein and immobilize it on the solid support.

[0136] Generally, an expressed T1R3 receptor can be used for HTS binding assays in conjunction with a ligand, such as an amino acid or carbohydrate. The identified peptide is labeled with a suitable radioisotope, including, but not limited to, <sup>125</sup>I, <sup>3</sup>H, <sup>35</sup>S or <sup>32</sup>P, by methods that are well known to those skilled in the art. Alternatively, the peptides may be labeled by well-known methods with a suitable fluorescent derivative (Baindur et al., Drug Dev. Res., 1994, 33, 373-398; Rogers, Drug Discovery Today, 1997, 2, 156-160). Radioactive ligand specifically bound to the receptor in membrane preparations made from the cell line expressing the recombinant protein can be detected in HTS assays in one of several standard ways, including filtration of the receptor-ligand complex to separate bound ligand from unbound ligand (Williams, Med. Res. Rev., 1991, 11, 147-184; Sweetnam et al., J. Natural Products, 1993, 56, 441-455). Alternative methods include a scintillation proximity assay (SPA) or a FlashPlate format in which such separation is unnecessary (Nakayama, Cur. Opinion Drug Disc. Dev., 1998, 1, 85-91; Bossé et al., J. Biomolecular Screening, 1998, 3, 285-292.). Binding of fluorescent ligands can be detected in various ways, including fluorescence energy transfer (FRET), direct spectrophotofluorometric analysis of bound ligand, or fluorescence polarization (Rogers, Drug Discovery Today, 1997, 2, 156-160; Hill, Cur. Opinion Drug Disc. Dev., 1998, I, 92-97).

[0137] Other assays may be used to identify specific ligands of a T1R3 receptor, including assays that identify ligands of the target protein through measuring direct binding of test ligands to the target protein, as well as assays that identify ligands of target proteins through affinity ultrafiltration with ion spray mass spectroscopy/HPLC methods or other physical and analytical methods. Alternatively, such binding interactions are evaluated indirectly using the yeast two-hybrid system described in Fields et al., Nature, 340:245-246 (1989), and Fields et al., Trends in Genetics, 10:286-292 (1994), both of which are incorporated herein by reference. The two-

hybrid system is a genetic assay for detecting interactions between two proteins or polypeptides. It can be used to identify proteins that bind to a known protein of interest, or to delineate domains or residues critical for an interaction. Variations on this methodology have been developed to clone genes that encode DNA binding proteins, to identify peptides that bind to a protein, and to screen for drugs. The two-hybrid system exploits the ability of a pair of interacting proteins to bring a transcription activation domain into close proximity with a DNA binding domain that binds to an upstream activation sequence (UAS) of a reporter gene, and is generally performed in yeast. The assay requires the construction of two hybrid genes encoding (1) a DNA-binding domain that is fused to a first protein and (2) an activation domain fused to a second protein. The DNA-binding domain targets the first hybrid protein to the UAS of the reporter gene; however, because most proteins lack an activation domain, this DNA-binding hybrid protein does not activate transcription of the reporter gene. The second hybrid protein, which contains the activation domain, cannot by itself activate expression of the reporter gene because it does not bind the UAS. However, when both hybrid proteins are present, the noncovalent interaction of the first and second proteins tethers the activation domain to the UAS, activating transcription of the reporter gene. For example, when the first protein is a receptor, or fragment thereof, that is known to interact with another protein or nucleic acid, this assay can be used to detect agents that interfere with the binding interaction. Expression of the reporter gene is monitored as different test agents are added to the system. The presence of an inhibitory agent results in lack of a reporter signal.

[0138] The yeast two-hybrid assay can also be used to identify proteins that bind to the gene product. In an assay to identify proteins that bind to a T1R3 receptor, or fragment thereof, a fusion polynucleotide encoding both a T1R3 receptor (or fragment) and a UAS binding domain (i.e., a first protein) may be used. In addition, a large number of hybrid genes each encoding a different second protein fused to an activation domain are produced and screened in the assay. Typically, the second protein is encoded by one or more members of a total cDNA or genomic DNA fusion library, with each second protein-coding region being fused to the activation domain. This system is applicable to a wide variety of proteins, and it is not necessary to know the identity or function of the second binding protein. The system is highly sensitive and can detect interactions not revealed by other methods; even transient interactions may trigger transcription to produce a stable mRNA that can be repeatedly translated to yield the reporter protein.

[0139] Other assays may be used to search for agents that bind to the target protein. One such screening method to identify direct binding of test ligands to a target protein is described in U.S. Patent No. 5,585,277, incorporated herein by reference. This method relies on the principle that proteins generally exist as a mixture of folded and unfolded states, and continually alternate between the two states. When a test ligand binds to the folded form of a target protein (i.e., when the test ligand is a ligand of the target protein), the target protein molecule bound by the ligand remains in its folded state. Thus, the folded target protein is present to a greater extent in the presence of a test ligand which binds the target protein, than in the absence of a ligand. Binding of the ligand to the target protein can be determined by any method that distinguishes between the folded and unfolded states of the target protein. The function of the target protein need not be known in order for this assay to be performed. Virtually any agent can be assessed by this method as a test ligand, including, but not limited to, metals, polypeptides, proteins, lipids, polysaccharides, polynucleotides and small organic molecules.

[0140] Another method for identifying ligands of a target protein is described in Wieboldt et al., Anal. Chem., 69:1683-1691 (1997), incorporated herein by reference. This technique screens combinatorial libraries of 20-30 agents at a time in solution phase for binding to the target protein. Agents that bind to the target protein are separated from other library components by simple membrane washing. The specifically selected molecules that are retained on the filter are subsequently liberated from the target protein and analyzed by HPLC and pneumatically assisted electrospray (ion spray) ionization mass spectroscopy. This procedure selects library components with the greatest affinity for the target protein, and is particularly useful for small molecule libraries.

[0141] Other embodiments of the invention comprise using competitive screening assays in which neutralizing antibodies capable of binding a polypeptide of the invention specifically compete with a test compound for binding to the polypeptide. In this manner, the antibodies can be used to detect the presence of any peptide that shares one or more antigenic determinants with T1R3 receptor. Radiolabeled competitive binding studies are described in A.H. Lin et al., Antimicrobial Agents and Chemotherapy, 1997, 41(10): 2127-2131, the disclosure of which is incorporated herein by reference in its entirety.

[0142] Another aspect of the present invention is directed to methods of identifying compounds that modulate (i.e., increase or decrease) activity of T1R3 comprising contacting T1R3 with a compound, and determining whether the compound modifies activity of T1R3. The activity in the presence of the test compound is compared to the activity in the absence of the test

compound. Where the activity of the sample containing the test compound is higher than the activity in the sample lacking the test compound, the compound is an agonist. Similarly, where the activity of the sample containing the test compound is lower than the activity in the sample lacking the test compound, the compound is an antagonist.

[0143] Agents that modulate (i.e., increase, decrease, or block) T1R3 activity or expression also may be identified, for example, by incubating a putative modulator with a cell containing a T1R3 polypeptide or polynucleotide and determining the effect of the putative modulator on T1R3 activity or expression. The selectivity of a compound that modulates the activity of T1R3 can be evaluated by comparing its effects on T1R3 to its effect on other T1R receptors. Selective modulators may include, for example, antibodies and other proteins, peptides, or organic molecules that specifically bind to a T1R3 polypeptide or a T1R3-encoding nucleic acid. Modulators of T1R3 activity will be therapeutically useful in treatment of diseases and physiological conditions in which normal or aberrant T1R3 activity is involved. Compounds identified as modulating T1R3 activity may be further tested in other assays including, but not limited to, in vivo models, in order to confirm or quantitate their activity.

[0144] The invention also provides methods for identifying a T1R3 modulator by: (a) contacting a T1R3 binding partner and a composition comprising a T1R3 receptor in the presence and in the absence of a putative modulator compound; (b) detecting binding between the binding partner and the T1R3 receptor; and (c) identifying a putative modulator compound or a modulator compound in view of decreased or increased binding between the binding partner and the T1R3 receptor in the presence of the putative modulator, as compared to binding in the absence of the putative modulator. Compounds identified as modulators of binding between T1R3 and a T1R3 binding partner may be further tested in other assays including, but not limited to, *in vivo* models, in order to confirm or quantitate their activity.

[0145] The invention also includes within its scope high-throughput screening (HTS) assays to identify compounds that interact with, enhance, or inhibit biological activity (i.e., affect enzymatic activity, binding activity, etc.) of a T1R3 polypeptide. HTS assays permit screening of large numbers of compounds in an efficient manner. Cell-based HTS systems are contemplated to investigate T1R3 receptor-ligand interaction. HTS assays are designed to identify "hits" or "lead compounds" having the desired property, from which modifications can be designed to improve the desired property. Chemical modification of the "hit" or "lead compound" is often based on an identifiable structure/activity relationship between the "hit" and the T1R3 polypeptide.

[0146] For example, modulators of T1R3 activity may be identified by expressing the T1R3 receptor in a heterologous cultured mammalian cell line, such as HEK cells, and detecting receptor activity in the presence and absence of a test compound by monitoring changes in intracellular calcium using a calcium-specific intracellular dye. In another embodiment, this process may be automated using a high-throughput screening device.

[0147] Candidate modulators contemplated by the invention include compounds selected from libraries of either potential activators or potential inhibitors. There are a number of different libraries used for the identification of small molecule modulators, including: (1) chemical libraries, (2) natural product libraries, and (3) combinatorial libraries comprised of random peptides, oligonucleotides, or organic molecules. Chemical libraries consist of random chemical structures, some of which are analogs of known compounds or analogs of compounds that have been identified as "hits" or "leads" in other drug discovery screens, some of which are derived from natural products, and some of which arise from non-directed synthetic organic chemistry. Natural product libraries are collections of microorganisms, animals, plants, or marine organisms that are used to create mixtures for screening by: (1) fermentation and extraction of broths from soil, plant, or marine microorganisms or (2) extraction of plants or marine organisms. Natural product libraries include polyketides, non-ribosomal peptides, and variants (non-naturally occurring) thereof. For a review, see Science 282:63-68 (1998). Combinatorial libraries are composed of large numbers of peptides, oligonucleotides, or organic compounds as a mixture. These libraries are relatively easy to prepare by traditional automated synthesis methods, PCR, cloning, or proprietary synthetic methods. Of particular interest are non-peptide combinatorial libraries. Still other libraries of interest include peptide, protein, peptidomimetic, multiparallel synthetic collection, recombinatorial, and polypeptide libraries. For a review of combinatorial chemistry and libraries created therefrom, see Myers, Curr. Opin. Biotechnol. 8:701-707 (1997). Identification of modulators through use of the various libraries described herein permits modification of the candidate "hit" (or "lead") to optimize the capacity of the "hit" to modulate activity.

[0148] T1R3 binding partners that stimulate T1R3 activity are useful as agonists in disease states or conditions characterized by insufficient T1R3 signaling (e.g., as a result of insufficient activity of a T1R3 ligand). T1R3 binding partners that block ligand-mediated T1R3 signaling are useful as T1R3 antagonists to treat disease states or conditions characterized by excessive T1R3 signaling. Thus, in another aspect, the invention provides methods for treating a disease or abnormal condition by administering to a patient in need of such treatment a substance that

modulates the activity or expression of a polypeptide having a sequence of SEQ ID NO:2, or exhibiting substantially the same biological activity as a polypeptide having a sequence of SEQ ID NO:2.

[0149] In addition T1R3 modulators in general, as well as T1R3 polynucleotides and polypeptides, are useful in diagnostic assays for such diseases or conditions.

#### **Mimetics**

[0150] Mimetics or mimics of compounds identified herein (sterically similar compounds formulated to mimic the key portions of the structure) may be designed for pharmaceutical use. Mimetics may be used in the same manner as the compounds identified by the present invention that modulate the T1R3 receptor and hence are also functional equivalents. The generation of a structural-functional equivalent may be achieved by the techniques of modeling and chemical design known to those of skill in the art. It will be understood that all such sterically similar constructs fall within the scope of the present invention.

[0151] The design of mimetics to a known pharmaceutically active compound is a known approach to the development of pharmaceuticals based on a "lead" compound. This is desirable where, for example, the active compound is difficult or expensive to synthesize, or where it is unsuitable for a particular method of administration, e.g., some peptides may be unsuitable active agents for oral compositions as they tend to be quickly degraded by proteases in the alimentary canal.

[0152] There are several steps commonly taken in the design of a mimetic. First, the particular parts of the compound that are critical and/or important in determining its T1R3-modulating properties are determined. In the case of a polypeptide, this can be done by systematically varying the amino acid residues in the peptide, e.g. by substituting each residue in turn. Alanine scans of peptides are commonly used to refine such peptide motifs.

[0153] Once the active region of the compound has been identified, its structure is modeled according to its physical properties, e.g. stereochemistry, bonding, size, and/or charge, using data from a range of sources, such as, but not limited to, spectroscopic techniques, X-ray diffraction data, and NMR. Computational analysis, similarity mapping (which models the charge and/or volume of the active region, rather than the bonding between atoms), and other techniques known to those of skill in the art can be used in this modeling process.

[0154] In a variant of this approach, the three-dimensional structure of the compound that modulates a T1R3 receptor and the active region of the T1R3 receptor are modeled. This can be especially useful where either or both of these compounds change conformation upon binding. Knowledge of the structure of the ligand-binding domain (for example, residues 1-571 of SEQ ID NO:2) of the receptor also allows the design of high potency ligands and/or modulators.

[0155] A template molecule is then selected onto which chemical groups that mimic the T1R3 modulator can be grafted. The template molecule and the chemical groups grafted onto it can conveniently be selected so that the mimetic is easy to synthesize, is pharmacologically acceptable, and does not degrade *in vivo*, while retaining the biological activity of the lead compound. Alternatively, where the mimetic is peptide-based, further stability can be achieved by cyclizing the peptide, thereby increasing its rigidity. The mimetic or mimetics found by this approach can then be screened by the methods of the present invention to see whether they have the ability to modulate the T1R3 receptor. Further optimization or modification can then be performed to arrive at one or more final mimetics for *in vivo* or clinical testing.

# Compositions of binding and/or modulating compounds

[0156] Following identification of a compound that binds and/or or modulates a T1R3 receptor, the compound may be manufactured and/or used in preparation of compositions including, but not limited to, foods, drinks, and pharmaceutical compositions. The compositions are provided or administered to patients, including, but not limited to, avians, felines, canines, bovines, ovines, porcines, equines, rodents, simians, and humans.

[0157] Thus, the present invention extends, in various aspects, not only to compounds identified in accordance with the methods disclosed herein but also foods, drinks, pharmaceutical compositions, drugs, or other compositions comprising such a compound; methods comprising administration of such a composition to a patient, e.g. for treatment (which includes prophylactic treatment) of a T1R3-associated disorder (e.g., obesity, diabetes); uses of such a compound in the manufacture of a composition for administration to a patient; and methods of making a composition comprising admixing such a compound with a pharmaceutically acceptable excipient, vehicle or carrier, and optionally other ingredients.

[0158] The compositions of the invention comprise a taste-modifying amount of at least one or more binding or modulating compounds. A "taste-modifying amount" is a quantity sufficient to increase or decrease the perception of a taste stimulus by a given mammal. The food and drink

compositions of the invention are formulated by the addition of a binding or modulating compound to a food or drink of the mammal. Such compositions may be individualized or breed-specific. For example, feline veterinary specialty diets may thus be made more palatable.

[0159] The pharmaceutical compositions of the invention comprise a therapeutically effective amount of a compound identified according to the methods disclosed herein, or a pharmaceutically acceptable salt thereof, and a pharmaceutically acceptable carrier or excipient.

[0160] The compounds of the invention can be formulated as neutral or salt forms. Pharmaceutically acceptable salts include those formed with free amino groups such as those derived from hydrochloric, phosphoric, acetic, oxalic, tartaric acids, etc., and those formed with free carboxyl groups such as those derived from sodium, potassium, ammonium, calcium, ferric hydroxides, isopropylamine, triethylamine, 2-ethylamino ethanol, histidine, procaine, etc.

[0161] Pharmaceutically acceptable carriers include but are not limited to saline, buffered saline, dextrose, water, glycerol, ethanol, and combinations thereof. The carrier and composition can be sterile. The formulation should suit the mode of administration.

[0162] The composition, if desired, can also contain minor amounts of wetting or emulsifying agents or pH buffering agents. The composition can be a liquid solution, suspension, emulsion, tablet, pill, capsule, sustained release formulation, or powder. The composition can be formulated as a suppository, with traditional binders and carriers such as triglycerides. Oral formulations can include standard carriers such as pharmaceutical grades of mannitol, lactose, starch, magnesium stearate, sodium saccharine, cellulose, magnesium carbonate, etc.

[0163] The pharmaceutical compositions of the invention may further comprise a secondary compound for the treatment of a disorder unrelated to the T1R3 receptor, such as an antibiotic or other therapeutic agent, to improve the palatability of the pharmaceutical composition, thereby improving the ease of administration.

[0164] In one embodiment, the composition is formulated in accordance with routine procedures as a pharmaceutical composition adapted for oral (e.g., tablets, granules, syrups) or non-oral (e.g., ointments, injections) administration to the subject. Various delivery systems are known and can be used to administer a compound that modulates a T1R3 receptor, e.g., encapsulation in liposomes, microparticles, microcapsules, expression by recombinant cells, receptor-mediated endocytosis, construction of a therapeutic nucleic acid as part of a retroviral or

other vector, etc. Methods of introduction include but are not limited to intradermal, intramuscular, intraperitoneal, intravenous, subcutaneous, intranasal, topical, and oral routes.

[0165] The compounds of the invention may be administered by any convenient route, for example by infusion or bolus injection, by absorption through epithelial or mucocutaneous linings (e.g., oral mucosa, rectal and intestinal mucosa, etc.), and may be administered together with other biologically active agents, for example in HAART therapy. Administration can be systemic or local. In addition, it may be desirable to introduce the pharmaceutical compositions of the invention into the central nervous system by any suitable route, including intraventricular and intrathecal injection; intraventricular injection may be facilitated by an intraventricular catheter, for example, attached to a reservoir, such as an Ommaya reservoir.

[0166] In a specific embodiment, it may be desirable to administer the pharmaceutical compositions of the invention locally to the area in need of treatment; this may be achieved by, for example, and not by way of limitation, local infusion during surgery; topical application, e.g., in conjunction with a wound dressing after surgery; by injection; by means of a catheter; by means of a suppository; or by means of an implant, said implant being of a porous, non-porous, or gelatinous material, including membranes, such as sialastic membranes, or fibers.

[0167] The composition can be administered in unit dosage form and may be prepared by any of the methods well known in the pharmaceutical art, for example, as described in REMINGTON'S PHARMACEUTICAL SCIENCES (Mack Publishing Co., Easton, PA). The amount of the compound of the invention that modulates a T1R3 receptor that is effective in the treatment of a particular disorder or condition will depend on factors including but not limited to the chemical characteristics of the compounds employed, the route of administration, the age, body weight, and symptoms of a patient, the nature of the disorder or condition, and can be determined by standard clinical techniques. Typically therapy is initiated at low levels of the compound and is increased until the desired therapeutic effect is achieved. In addition, in vitro assays may optionally be employed to help identify optimal dosage ranges. Suitable dosage ranges for intravenous administration are preferably generally about 20-500 micrograms of active compound per kilogram body weight. Suitable dosage ranges for intranasal administration are preferably generally about 0.01 pg/kg body weight to 1 mg/kg body weight. Suppositories preferably generally contain active ingredient in the range of 0.5% to 10% by weight; oral formulations preferably may contain 10% to 95% active ingredient. Effective doses may be extrapolated from dose-response curves derived from in vitro or animal model test systems.

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[0168] Typically, compositions for intravenous administration are solutions in sterile isotonic aqueous buffer. Where necessary, the composition may also include a solubilizing agent and a local anesthetic such as lidocaine to ease pain at the site of the injection. Generally, the ingredients are supplied either separately or mixed together in unit dosage form, for example, as a dry-lyophilized powder or water-free concentrate in a hermetically sealed container such as an ampoule or sachette indicating the quantity of active agent. Where the composition is to be administered by infusion, it can be dispensed with an infusion bottle containing sterile pharmaceutical grade water or saline.

[0169] Where the composition is administered by injection, an ampoule of sterile water for injection or saline can be provided so that the ingredients may be mixed prior to administration.

### **Treatment Methods**

[0170] The invention provides methods of treatment of T1R3 receptor-associated disorders by administering to a subject or patient an effective amount of a compound that modulates the T1R3 receptor. In some aspects of the invention, the compounds or pharmaceutical compositions of the invention are administered to a patient having an increased risk of or having a disorder associated with the T1R3 receptor. The patient may be, for example, avian, feline, canine, bovine, ovine, porcine, equine, rodent, simian, or human.

## Kits

[0171] A kit of the invention comprises a carrier means being compartmentalized to receive in close confinement one or more container means such as vials, tubes, and the like, each of the container means comprising an element to be used in the methods of the invention. For example, one of the container means may comprise the a polynucleotide encoding a T1R3 receptor of the invention, a T1R3 receptor of the invention, or an antibody thereto. The kit may also have one or more conventional kit components, including, but not limited to, instructions, test tubes, Eppendorf<sup>TM</sup> tubes, labels, reagents helpful for quantification of marker gene expression, *etc*.

## **EXAMPLES**

[0172] The following examples are meant to be illustrative of the present invention and are not intended to limit the scope thereof.

#### Cloning and Characterization of the Feline T1R3 receptor

[0173] The discovery of feline taste receptor, T1R3, was achieved by using a molecular strategy termed "overgo" (Thomas, et al., Genome Res., 12:1277-1285 (2002); Vollrath, D., DNA markers for physical mapping In GENOME ANALYSIS: A LABORATORY MANUAL, Vol. 4, ed. B. Birren, et al., pp. 187-215, 1999). Cold Spring Harbor Laboratory Press, Cold Spring Harbor, New York.). This strategy involves the use of the shortest DNA probes among the many kinds of probes used in bacterial artificial chromosome (BAC) library screening. These probes are comprised of two DNA sequences (e.g., 22mers or 24mers) with a complementary 8 base overlap. They can be designed by computer program (genome.wustl.edu/tools/?overgo=1) and are readily synthesized.

[0174] Overgo probes were designed from conserved regions of the chromosome 1 marker, "disheveled 1" (DVL1) and the G protein-coupled receptor, T1R3, by aligning DVL1 and T1R3 genomic sequences from many different species. The overlapping sequences of the seven DVL1 overgo probes used in the present invention were as follows:

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catoV1a ACTTTGAGAACATGAGTAATGACG (SEQ ID NO:21) catoV1b AGTACCCGGACTGCGTCGTCATTA (SEQ ID NO:22) catoV2a CACTAGGGTCATCCTTGCTTTCAG (SEQ ID NO:23) catoV2b AGTCAGGGTGATGGGCCTGAAAGC (SEQ ID NO:24) Ov8-OVa ATGTGGTGGACTGGCTGTACCATC (SEQ ID NO:25) Ov8-OVb TTGAAGCCCTCCACGTGATGGTAC (SEQ ID NO:26) Ov9a CACACGGTGAACAAGATCACCTTC (SEQ ID NO:27) Ov9b AGTAGCACTGCTCGGAGAAGGTGA (SEQ ID NO:28) Ov10a ATCTACCACATGGACGAGGAGGAG (SEQ ID NO:29) Ov10b TGACCAGGTACGGCTTCCTCCT (SEQ ID NO:30) Ov11a AGCGCGTCACGCTGGCCGACTTCA (SEQ ID NO:31) Ov11b TTGCTGAGCACGTTCTTGAAGTCG (SEQ ID NO:32) Ov12a CACGCCTACAAATTCTTCTTTAAG (SEQ ID NO:33) Ov12b AGTCCTGGTCCATGGACTTAAAGA (SEQ ID NO:34).
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The overlapping sequences of the twelve T1R3 overgo probes used in the present invention were as follows:

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tlr3-OVla CTTCCACTCCTGCTGCTACGACTG (SEQ ID NO:35)
tlr3-OVlb TGCCTCGCAGTCCACGCAGTCGTA (SEQ ID NO:36)
tlr3-OV2a AGGTGCGCCGCGTCAAGGGCTTCC (SEQ ID NO:37)
tlr3-OV2b TCGTAGCAGCAGGAGTGGAAGCCC (SEQ ID NO:38)
tlr3-OV3a GTTCCTGGCATGGGGGGAGCCGGC (SEQ ID NO:39)
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tlr3-0V3b GAGCAGCACAAGCACAGCCGGCTC (SEQ ID NO:40)
t1r3-OV4a ACAGCCCACTAGTTCAGGCCGCAG (SEQ ID NO:41)
t1r3-OV4b CAGGCCCGGGGTCCCCCTGCGGCC (SEQ ID NO:42)
t1r3-OV5a CCCACTGGTTCAGGCCTCGGGGGG (SEQ ID NO:43)
t1r3-oV5b AAAGCAGGCCAGGGGCCCCCCGA (SEQ ID NO:44)
t1r3-oV6a AGGCGCTGGTGCACTGCCGCACAC (SEQ ID NO:45)
t1r3-oV6b AAGCTGACCCAGGAGCGTGTGCGG (SEQ ID NO:46)
t1r3-OV7a ACAGAGGCACTGGTGCACTGCCGC (SEQ ID NO:47)
tlr3-0V7b TGATCCAGGAGTGCACGCGGCAGT (SEQ ID NO:48)
t1r3-OV8a ACCAATGCCACGCTGGCCTTTCTC (SEQ ID NO:49)
t1r3-ov8b AAGTGCCCAGGAAGCAGAGAAAGG (SEQ ID NO:50)
t1r3-OV9a TGGTACATGCTGCCAATGCCACGC (SEQ ID NO:51)
tlr3-OV9b AAGCAGAGGAAAGCCAGCGTGGCA (SEQ ID NO:52)
t1r3-OV10a TACAACCGTGCCCGTGGCCTCACC (SEQ ID NO:53)
t1r3-OV10b AGGCCAGCATGGCGAAGGTGAGGC (SEQ ID NO:54)
t1r3-OV11a TCATCACCTGGGTCTCCTTTGTGC (SEO ID NO:55)
t1r3-oV11b ACATTGGCCAGGAGGGGCACAAAG (SEQ ID NO:56)
t1r3-OV12a TGCAGATGGGTGCCCTCCTGCTCT (SEQ ID NO:57)
t1r3-OV12b AGGATGCCCAGCACACAGAGCAGG (SEQ ID NO:58).
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The single-stranded overhangs were filled in with <sup>32</sup>P labeled dATP and dCTP, and the overgo probes hybridized with BAC libraries.

[0175] The overgo strategy is considered to be more versatile than a PCR-based strategy by those skilled in the art of comparative physical mapping for the following reasons: (1) overgo probes are short (e.g., 36mers or 40mers), making the probability of good alignment from among many species more favorable; (2) overgo probes are more specific to the target genes compared with traditional cDNA and genomic DNA probes used by PCR; and (3) although overgo probes are short, they are not as restricted as traditional PCR probes, which cannot tolerate even a few mismatches, because they can be used in hybridization approaches with BACs or other libraries.

[0176] Screening a feline genomic BAC library. Seven DVL1 overgo probes (SEQ ID NOS:21-34) were used in screening a feline genomic BAC library. Probes were radioactively labeled by the random hexa-nucleotide method (Feinberg & Vogelstein, *Analytical Biochemistry*, 132:6-13 (1983)). Hybridization and washing of membranes followed standard protocols (Church & Gilbert, *PNAS U.S.A.*, 81:1991-1995 (1984)). Thirty-nine positive BAC clones were

identified. Several BAC ends were sequenced. One clone containing homologous sequence to human chromosome 1p36, BAC 552J19, was identified using bioinformatics tools.

[0177] Production of a shotgun library for BAC 552J19 and identification of a single clone containing feline T1R3. BAC DNA from 552J19 was prepared by using Qiagen Large Construct Kit. DNA was then digested by the restriction enzyme Sau3A1 and subcloned into pGEM+3Z (Promega) vector. After transformants were arrayed to a nylon membrane, two separate hybridizations were performed using seven DVL1 and twelve T1R3 overgo probes (SEQ ID NOS:35-58). Two clones positive for DVL1 and four clones positive for T1R3 were found. These clones were confirmed by sequencing. Because DVL1 is the neighboring gene of T1R3 in human and mouse, it is likely this also is the case in cat; therefore, the DVL1 positive clones verified that the BAC 552J19 is the correct BAC, that is, it is the one containing feline T1R3.

#### Results

[0178] More than 3 kb of genomic sequences containing the open reading frame for domestic cat taste receptor, T1R3, were obtained. Figure 1 shows the multiple sequence alignments of the known nucleotide sequences for the T1R receptors human (T1R1, SEQ ID NO:8; T1R2, SEQ ID NO:5; T1R3, SEQ ID NO:11), mouse (T1R1, SEQ ID NO:6; T1R2, SEQ ID NO:3; T1R3, SEQ ID NO:9), and rat (T1R1, SEQ ID NO:7; T1R2, SEQ ID NO:4; T1R3, SEQ ID NO:10), along with the newly discovered and novel nucleotide sequence for the T1R3 taste receptor from domestic cat (SEQ ID NO:1).

[0179] Figure 2 shows the deduced amino acid sequence of the domestic cat taste receptor, T1R3 (SEQ ID NO:2), aligned with the amino acid sequences of the T1R receptor family human (T1R1, SEQ ID NO:17; T1R2, SEQ ID NO:20; T1R3, SEQ ID NO:12), rat (T1R1, SEQ ID NO:16; T1R2, SEQ ID NO:19; T1R3, SEQ ID NO:14), and mouse (T1R1, SEQ ID NO:15; T1R2, SEQ ID NO:18; T1R3, SEQ ID NO:13). The deduced cat sequence predicts four additional amino acids at positions 11 - 14 relative to the homologous T1R3 receptors of mouse, human, and rat. The deduced sequence for cat reveals a threonine in position 64, a position equivalent to amino acid 60 in mouse, and a leucine at position 59, a position equivalent to position 55 in mouse. In mouse, amino acid substitutions of a threonine at position 60 and an alanine at position 55, both positions located within the putative extracellular N-terminal domain of the polypeptide, are present in strains of mice demonstrating low preference for the sweet stimulus saccharin (Bachmanov *et al.*, *Chem. Senses*, 26:925-933 (2001)). Leucine is a

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conservative substitution for alanine. Accordingly, the amino acid sequence differences of cat and mouse T1R3 receptor may account for functional differences that lead to different taste preferences between the two species. For example, the amino acid substitutions may explain the cat's inability to taste many compounds that have a sweet taste to mice and humans.

[0180] Table 4 shows the percent homology among the members of the T1R family in relation to the newly discovered cat T1R3 taste receptor. The portion of Table 4 to the left of the diagonal (in bold type) shows the percent homology based on the open reading frame of the nucleotide sequences obtained from Figure 1 for the T1R family among human, cat, rat, and mouse. The upper portion to the right of the diagonal (in italic type) shows the percent homology of the T1R members based on the amino acid sequences of Figure 2. Cat T1R3 shows 79% nucleotide sequence homology with human T1R3, 75% with rat T1R3 and 74% with mouse T1R3. At the amino acid level, cat T1R3 shows 73% homology with human T1R3, 72% with rat, and 72% with mouse. Cat T1R3 shows generally low homology with the other known members of the T1R family, T1R1 and T1R2, from human, rat, and mouse. The same range of relatively low homology is present among the human, rat, and mouse T1R3 and the T1R1 and T1R2 receptors from the same species.

Table 4. Percent Homology Among Diverse Species for T1Rs

Species	Mouse T1R1	Mouse T1R2	Mouse T1R3	Rat T1R1	Rat T1R2	Rat T1R3	Human T1R1	Human T1R2	Human T1R3	Cat T1R3
Mouse T1R1		36	30	90	36	30	73	37	30	30
Mouse T1R2	55		28	36	91	28	34	69	28	28
Mouse T1R3	33	15		31	28	92	30	27	72	72
Rat T1R1	91	55	33		37	31	73	37	31	31
Rat T1R2	55	91	15	57		28	34	71	29	28
Rat T1R3	33	21	93	32	15		31	27	73	72
Human T1R1	79	56	35	79	56	35		35	31	31
Human T1R2	57	78	17	56	78	17	57		28	28
Human T1R3	41	39	73	39	36	75	40	38		73
Cat T1R3	33	34	74	36	36	75	53	39	79	

Note: Upper right cells (italics) contain deduced amino acid homology; lower left cells (bold) contain nucleotide homology.

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[0181] The rat and mouse have closely related T1R receptors, while the T1R3 of human and cat diverge from these two, as illustrated in the phylogenetic tree of Figure 3. Interestingly, the types of sweet compounds to which the rat and mouse respond are very similar, whereas those that stimulate the human and those that stimulate the cat are much different from those for rat and mouse, and whereas the compounds that stimulate the cat and human receptors also are very different.

[0182] The feline T1R3 receptor is a seven transmembrane receptor similar in structure to other known members of the T1R family of receptors (Figure 4). The structure of the feline T1R3 receptor was generated through use of a protein modeling program available at <a href="https://www.ebi.ac.uk/~moeller/transmembrane.html">www.ebi.ac.uk/~moeller/transmembrane.html</a>.

#### What is Claimed:

- 1. An isolated and purified polynucleotide encoding a T1R3 receptor comprising:
  - a) the nucleotide sequence of SEQ ID NO:1,
- b) a fragment of at least about 42 contiguous nucleotides of SEQ ID NO:1 encoding a polypeptide having substantially the same biological activity as a polypeptide encoded by the nucleotide sequence of SEQ ID NO:1,
- c) a variant of the polynucleotide of SEQ ID NO:1 having at least 80% homology to the polynucleotide of SEQ ID NO:1 and encoding a polypeptide having substantially the same biological activity as a polypeptide encoded by the nucleotide sequence of SEO ID NO:1,
- d) a variant of the polynucleotide of SEQ ID NO:1 having at least 80% homology to the polynucleotide of SEQ ID NO:1 and encoding a polypeptide conferring modified taste perception to one or more taste stimuli relative to a polypeptide encoded by the polynucleotide of SEQ ID NO:1,
  - e) a nucleotide sequence encoding the amino acid sequence of SEQ ID NO:2,
- f) a nucleotide sequence substantially complementary to the nucleotide sequence of SEQ ID NO:1, or
- g) a nucleotide sequence that hybridizes to the complement of the polynucleotide having SEQ ID NO:1 under high stringency conditions.
- 2. The polynucleotide of claim 1, wherein said polynucleotide is DNA.
- 3. The polynucleotide of claim 1, wherein said polynucleotide is RNA.
- 4. The polynucleotide of claim 1 comprising a variant of the polynucleotide of SEQ ID NO:1 encoding an amino acid sequence of SEQ ID NO:2 having a nonconserved amino acid substitution at residue 59 or residue 64.
- 5. The polynucleotide of claim 1 comprising a fragment of the polynucleotide of SEQ ID NO:1 wherein said fragment comprises a nucleotide sequence encoding an extracellular domain of the polypeptide of SEQ ID NO:2, a transmembrane domain of the polypeptide of SEQ ID NO:2, or an intracellular domain of the polypeptide of SEQ ID NO:2.

- 6. An expression vector comprising the polynucleotide of claim 1 operably linked to a promoter.
- 7. A host cell comprising the expression vector of claim 6.
- 8. The host cell of claim 7 wherein said cell is mammalian.
- 9. The host cell of claim 8 wherein said cell is feline.
- 10. A cell culture comprising at least one cell of claim 6.
- 11. An isolated and purified T1R3 receptor polypeptide encoded by a polynucleotide of claim 1.
- 12. The polypeptide of claim 11 wherein said polypeptide comprises the amino acid sequence of SEQ ID NO:2, a fragment of at least 30 contiguous amino acids of SEQ ID NO:2, or a variant thereof having substantially the same biological activity as the polypeptide of SEQ ID NO:2.
- 13. The polypeptide of claim 11 wherein said polypeptide comprises an amino acid sequence having at least one sequence variation of SEQ ID NO:2 wherein said variation confers modified taste perception to one or more taste stimuli relative to a polypeptide of SEQ ID NO:2.
- 14. An isolated and purified T1R3 receptor polypeptide comprising the amino acid sequence of SEQ ID NO:2.
- 15. The polypeptide of claim 12, wherein said polypeptide comprises a feline T1R3 receptor.
- 16. A kit for the detection of a polynucleotide encoding a feline T1R3 receptor comprising a polynucleotide that specifically hybridizes to a polynucleotide encoding a polypeptide having an amino acid sequence of SEQ ID NO:2 and instructions relating to detection of said polynucleotide.
- 17. An isolated and purified antibody that immunoreacts specifically with at least one epitope of a polypeptide of one of claims 11, 12, 13, 14, or 15.
- 18. A kit for the detection of a polypeptide encoding a feline T1R3 receptor comprising the antibody of claim 17 and instructions relating to detection.
- 19. A method of producing a feline T1R3 receptor comprising culturing the host cell of claim 7 and recovering said receptor from said host cell.

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- 20. The feline T1R3 receptor produced according to the method of claim 19.
- 21. A method for identifying compounds that interact with a feline T1R3 receptor comprising:

contacting a feline T1R3 receptor of claim 11, 12, 13, 14, or 15 with a test compound, and

detecting interaction between said receptor and said compound.

- 22. The method of claim 21, wherein said receptor is bound to a solid support.
- 23. The method of claim 22, wherein said solid support is formulated into a feline-specific electronic tongue.

A method for identifying an agonist of a feline T1R3 receptor comprising:

- expressing an expression vector of claim 6 in the presence of a test compound, and detecting an increase in biological activity of a polypeptide produced by said expression step in the presence of said compound relative to biological activity of said polypeptide in the absence of said compound.
- 25. A method for identifying an agonist of a feline T1R3 receptor comprising:
  contacting a polypeptide of claim 11, 12, 13, 14, or 15 with a test compound, and
  detecting an increase in biological activity of said polypeptide in the presence of said
  compound relative to biological activity of said polypeptide in the absence of said compound.
- 26. A method for identifying an antagonist of a feline T1R3 receptor comprising:
  expressing an expression vector of claim 6 in the presence of a test compound, and
  detecting a decrease in biological activity of a polypeptide produced by said expression
  step in the presence of said compound relative to biological activity of said polypeptide in the absence of said compound.
- 27. A method for identifying an antagonist of a feline T1R3 receptor comprising: contacting the polypeptide of claim 11, 12, 13, 14, or 15 with a test compound, and

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detecting a decrease in biological activity of said polypeptide in the presence of said compound relative to biological activity of said polypeptide in the absence of said compound.

- 28. A compound for modifying the taste perception of a mammal identified according to the method of claim 24.
- 29. A compound for modifying the taste perception of a mammal identified according to the method of claim 25.
- 30. A compound for modifying the taste perception of a mammal identified according to the method of claim 26.
- 31. A compound for modifying the taste perception of a mammal identified according to the method of claim 27.
- 32. A composition for modifying the taste perception of a mammal comprising at least one compound identified by the method of claim 24.
- 33. A composition for modifying the taste perception of a mammal comprising at least one compound identified by the method of claim 25.
- 34. A composition for modifying the taste perception of a mammal comprising at least one compound identified by the method of claim 26.
- 35. A composition for modifying the taste perception of a mammal comprising at least one compound identified by the method of claim 27.
- 36. The composition of claim 32, wherein said composition comprises a feline food, drink, or pharmaceutical composition.
- 37. The composition of claim 33, wherein said composition comprises a feline food, drink, or pharmaceutical composition.
- 38. The composition of claim 34, wherein said composition comprises a feline food, drink, or pharmaceutical composition.
- 39. The composition of claim 35, wherein said composition comprises a feline food, drink, or pharmaceutical composition.

- 40. The composition of claim 36, wherein said composition is breed-specific.
- 41. The composition of claim 37, wherein said composition is breed-specific.
- 42. The composition of claim 38, wherein said composition is breed-specific.
- 43. The composition of claim 39, wherein said composition is breed-specific.
- 44. A method for modifying the taste perception of a mammal comprising administering to said mammal a compound identified according to the method of claim 24.
- 45. A method for modifying the taste perception of a mammal comprising administering to said mammal a compound identified according to the method of claim 25.
- 46. A method for modifying the taste perception of a mammal comprising administering to said mammal a compound identified according to the method of claim 26.
- 47. A method for modifying the taste perception of a mammal comprising administering to said mammal a compound identified according to the method of claim 27.
- 48. A composition for modifying the taste perception of a mammal comprising at least one polynucleotide of claim 1 and a pharmaceutically acceptable excipient.
- 49. A method for modifying the taste perception of a mammal comprising administering to said mammal at least one polynucleotide of claim 1.
- 50. A method for modifying the taste perception of a mammal comprising administering to said mammal at least one polypeptide of claim 11.
- 51. A transgenic animal comprising a polynucleotide of claim 1.
- 52. The method of claim 25 wherein said polypeptide is bound to a solid support.
- 53. The method of claim 52 wherein said solid support is formulated into a feline-specific electronic tongue.
- 54. The method of claim 27 wherein said polypeptide is bound to a solid support.
- 55. The method of claim 54 wherein said solid support is formulated into a feline-specific electronic tongue.

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56. A method of identifying a feline T1R3 receptor variant that confers modified taste perception comprising expressing a variant of the polynucleotide of SEQ ID NO:1 having at least 80% homology to the polynucleotide of SEQ ID NO:1 and detecting an increase or a decrease in the biological activity of the polypeptide encoded by the variant relative to the biological activity of the polypeptide encoded by SEQ ID NO:1.

## **ABSTRACT**

The present invention relates to the discovery of a gene of the domestic cat (*Felis catus*) associated with taste perception. The invention provides, *inter alia*, the nucleotide sequence of the feline T1R3 receptor gene, the amino acid sequence of the polypeptide encoded thereby, and antibodies to the polypeptide. The present invention also relates to methods for screening for compounds that modify the gene's function or activity, the compounds identified by such screens, and mimetics of the identified compounds. The invention further provides methods for modifying the taste preferences, ingestive responses, or general behavior of a mammal by administering compounds that affect the function or activity of the gene.

Docket No.: MON-3042
App No.: Not Yet Assigned
Title: TASTE RECEPTOR OF THE T1R FAMILY FROM DOMESTIC CAT
Inventors: Xia L1, Weihua L1, Danielle R. REED, Alexander Alexeyevich BACHMANOV,
Joseph G. BRAND
Attomey: Felicity E. Groth
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### Figure 1A CLUSTAL W (1.82) multiple nucle tide sequence alignment of T1Rs:

mouseTas1r2 ratTas1r2 humanTAS1R2 mouseTas1r1 ratTas1r1 humanTAS1R1 mouseTas1r3 ratTas1r3 catTas1r3 humanTAS1R3		52 52 59 53 56 45 45
mouseTas1r2	TGCCTAAGCCAGTCATGCTGGTAGGGAAC-TCCGACTTTCACCTGGCTGGGGACTAC	108
ratTas1r2	TGCCTAAGCCAGGCAAGCTGGTAGAGAAC-TCTGACTTCCACCTGGCCGGGGACTAC	108
humanTAS1R2		99
mouseTaslrl	GGCTTTCAGCTGCCAAAG-GACAGAATCC-TCTCCAGGTTTCAGCCTCCCTGGGGACTTC	117
ratTas1r1	GGCTTTCAGCTGCCAAAG-GACAGAGTCC-TCTCCAGGCTTCAGCCTTCCTGGGGACTTC	
humanTAS1R1		114
mouseTaslr3		105
ratTas1r3	GAGCTTGGGATGGGGTCCTCTTTGTGTCTCTCACAGCAATTCAAGGCACAAGGGGACTAT	
catTas1r3	GACCACGGGGAGGCGCAACGTCCTGCTTGTCACAGCAGCTCAGGATGCAGGGGGACTAT	
humanTAS1R3	CACCCTGGGACGGGGCCCCATTGTGCCTGTCACAGCAACTTAGGATGAAGGGGGACTAC	105
	** * ** *	
mouseTaslr2	CTCCTGGGTGGCCTCTTTACCCTCCATGCCAACGTGAAGAGCGTCTCTCACCTCAGCTAC	168
ratTas1r2	CTCCTGGGTGGCCTCTTTACCCTCCATGCCAACGTGAAGAGCATCTCCCACCTCAGCTAC	
humanTAS1R2	CTCCTGGGTGGCCTCTTCTCCCTCCATGCCAACATGAAGGGCATTGTTCACCTTAACTTC	
mouseTaslr1	CTCCTGGCAGGCCTGTTCTCCCTCCATGCTGACTGTCTGCAGGTGAGACACAGAC	
ratTas1r1	CTCCTTGCAGGTCTGTTCTCCCTCCATGGTGACTGTCTGCAGGTGAGACACAGAC	
humanTAS1R1	CTCCTGGCAGGCCTGTTCCCTCTCCATTCTGGCTGTCTGCAGGTGAGGCACAGAC	
mouseTas1r3	ATACTGGGCGGGCTATTTCCCCTGGGCTCAACCGAGGAGGCCACTCTCAACCAG	159
ratTas1r3	ATATTGGGTGGACTATTTCCCCTGGGCACAACTGAGGAGGCCACTCTCAACCAG	
catTas1r3	GTGCTGGGTGGGCTCTTCCCTCTGGGCTCTGCCGAGGGTACAGGTCTTGGCGAC	171
humanTAS1R3	GTGCTGGGGGGGCTGTTCCCCCTGGGCGAGGCCGAGGAGGCTGGCCTCCGCAGC	159
	* * * ** ** * * * * *	
mouseman1 x2	CTGCAGGTGCCCAAGTGCAATGAGTACAACATGAAGGTCTTGGGCTACAACCTCATG	225
mouseTaslr2 ratTaslr2	CTGCAGGTGCCCAAGTGCAATGAGTACAACATGAAGGTGTTGGGCTACAACCTCATG	
humanTAS1R2		216
mouseTaslrl		231
ratTas1r1	CT-CTGGTGACAAGTTGTGACAGGCCCGACAGCTTCAACGGCCATGGCTACCACCTCTTC	
humanTAS1R1		228
mouseTaslr3		219
ratTas1r3	AGAACACAGCCCAACGGCATCCTATGTACCAGGTTCTCGCCCCTTGGTTTGTTCCTGGCC	
catTas1r3	GGGCTGCAGCCCAATGCCACCGTGTGCACCAGGTTCTCGTCTCTGGGCCTGCTCTGGGCG	
humanTAS1R3	CGGACACGGCCCAGCAGCCCTGTGTGCACCAGGTTCTCCTCAAACGGCCTGCTCTGGGCA	
	* * ** *	
	03.00003.00003.00003.0003.0003.0003.00	0.05
mouseTas1r2	CAGGCCATGCGATTCGCCGTGGAGGAAATCAACAACTGTAGCTCTCTGCTGCCCGGCGTG	
ratTas1r2	CAGGCCATGCGTTTCGCTGTGGAGGAGATCAACAACTGTAGCTCCCTGCTACCCGGCGTG	
humanTAS1R2	CAGGCCATGCGCTTCGCGGTGGAGGAGATCAACAATGACAGCAGCCTGCTGCCTGGTGTG CAAGCCATGCGGTTCACCGTTGAGGAGATAAACAACTCCACAGCTCTGCTTCCCAACATC	276
mouseTaslrl		285
ratTas1r1 humanTAS1R1		288
mouseTas1r3		279
ratTas1r3	ATGGCTATGAAGATGGCTGTAGAGGAGATCAACAATGGATCTGCCTTGCTCCCTGGGCTG  ATGGCTATGAAGATGGCTGTAGAGGAGATCAACAATGGATCTGCCTTGCTCCCTGGGCTG	
catTas1r3	CTGGCCGTGAAGATGGCGTGGAGGAGATCAACAATGGATCTGCCTTGCTCCTGGGCTG	
humanTAS1R3	CTGGCCGTGAAAATGGCCGTGGAGGAGATCAACAACAAGTCGGATCTGCTGCCCGGGCTG	
	** **	

Docket No.. MON-3042
App No.: Not Yet Assigned Filed: Herewith
Title: TASTE RECEPTOR OF THE TIR FAMILY FROM DOMESTIC CAT
Inventors: Xia Li, Weihua Ll, Danielle R. REED, Alexander Alexeyevich BACHMANOV,
Joseph G. BRAND
Attorney: Felicity E. Groth Phone: (215) 568-3100
Sheet 2 of 13

# Figure 1B

mouseTas1r2	CTGCTCGGCTACGAGATGGTGGATGTCTGCTACCTCTCCAACAATATCCAGCCTGGG	342
ratTas1r2	CTGCTCGGCTACGAGATGGTGGATGTCTGTTACCTCTCCAACAATATCCACCCTGGG	342
humanTAS1R2	CTGCTGGGCTATGAGATCGTGGATGTGTGCTACATCTCCAACAATGTCCAGCCGGTG	333
mouseTaslrl	ACCCTGGGGTATGAACTGTATGACGTGTGCTCAGAGTCTTCCAATGTCTATGCCACC	348
ratTaslrl	ACCCTGGGGTATGAGCTGTACGACGTGTGCTCAGAATCTGCCAATGTGTATGCCACC	342
humanTAS1R1	ACCCTGGGGTACCAGCTGTATGATGTGTGTTCTGACTCTGCCAATGTGTATGCCACG	345
mouseTas1r3	CGGCTGGGCTATGACCTATTTGACACATGCTCCGAGCCAGTGGTCACCATGAAATCCAGT	339
ratTaslr3	CGACTGGGCTATGACCTGTTTGACACATGCTCAGAGCCAGTGGTCACCATGAAGCCCAGC	339
catTaslr3	CACCTGGGCTATGACCTCTTTGACACGTGTTCAGAGCCCATGGTGGCCATGAAGCCCAGC	351
humanTAS1R3	CGCCTGGGCTACGACCTCTTTGATACGTGCTCGGAGCCTGTGGTGGCCATGAAGCCCAGC	339
	** ** ** * * * * * * * * * * * * * * * *	
mouseTas1r2	CTCTACTTCCTGTCACAGATAGATGACTTCCTGCCCATCCTCAAAGACTACAGCCAG	399
ratTaslr2	CTCTACTTCCTGGCACAGGACGACGTCCTGCCCATCCTCAAAGACTACAGCCAG	399
humanTAS1R2	CTCTACTTCCTGGCACACGAGGACAACCTCCTTCCCATCCAAGAGGACTACAGTAAC	390
mouseTas1r1	CTGAGGGTGCTCGCCCAGCAAGGGACAGGCCACCTAGAGATGCAGAGAGATCTTCGCAAC	408
ratTas1r1	CTGAGGGTGCTCGCCCGCAAGGGCCCCGCCACATAGAGATACAGAAAGACCTTCGCAAC	402
humanTASIR1	CTGAGAGTGCTCTCCCTGCCAGGGCAACACCACATAGAGCTCCAAGGAGACCTTCTCCAC	405
mouseTas1r3	CTCATGTTCCTGGCCAAGGTGGGCAGTCAAAGCATTGCTGCCTACTGCAACTACACACAG	399
ratTaslr3	CTCATGTTCATGGCCAAGGTGGGAAGTCAAAGCATTGCTGCCTACTGCAACTACACACAG	399
catTaslr3	CTCGTGTTCATGGCCAAAGCAGGCAGCTGCAGCATTGCCGCCTACTGCAATTACACACAG	411
humanTAS1R3	CTCATGTTCCTGGCCAAGGCAGGCAGCCGCGACATCGCCGCCTACTGCAACTACACGCAG	399
	** * * * * * * * *	
mouseTas1r2	TACAGGCCCCAAGTGGTGGCCGTCATTGGCCCAGACACTCTGAGTCCGCCATCACCGTG	459
ratTas1r2	TACATGCCCCACGTGGTGGCTGTCATTGGCCCCGACAACTCTGAGTCCGCCATTACCGTG	459
humanTAS1R2	TACATTTCCCGTGTGGTGGCTGTCATTGGCCCTGACAACTCCGAGTCTGTCATGACTGTG	450
mouseTas1r1	CACTCCTCCAAGGTGGTGGCACTCATTGGGCCTGATAACACTGACCACGCTGTCACCACT	468
ratTaslrl	CACTCCTCCAAGGTGGTGGCCTTCATCGGGCCTGACAACACTGACCACGCTGTCACTACC	462
humanTAS1R1	TATTCCCCTACGGTGCTGGCAGTGATTGGGCCTGACAGCACCAACCGTGCTGCCACCACA	465
mouseTaslr3	TACCAACCCCGTGTGCTGGCTGTCATCGGCCCCCACTCATCAGAGCTTGCCCTCATTACA	459
ratTas1r3	TACCAACCCCGTGTGCTGGCTGTCATTGGTCCCCACTCATCAGAGCTTGCCCTCATTACA	459
catTas1r3	TACCAGCCCGCGTGCTGGCCGTCATCGGGCCCCACTCGTCTGAGCTCGCCCTCGTCACC	471
humanTAS1R3	TACCAGCCCGTGTGCTGGCTGTCATCGGGCCCCACTCGTCAGAGCTCGCCATGGTCACC	459
mouseTas1r2	TCCAACATTCTCTCCTACTTCCTCGTGCCACAGGTCACATATAGCGCCATCACCGACAAG	519
ratTas1r2	TCCAACATTCTCTCATTTCCTCATCCCACAGATCACATACAGCGCCATCTCCGACAAG	519
humanTAS1R2	GCCAACTTCCTCTCCCTATTTCTCCTTCCACAGATCACCTACAGCGCCATCAGCGATGAG	510
mouseTaslr1	GCTGCCCTGCTGAGCCCTTTTCTGATGCCCCTGGTCAGCTATGAGGCGAGCAGCGTGATC	528
ratTas1rl	GCTGCCTTGCTGGGTCCTTTCCTGATGCCCCTGGTCAGCTATGAGGCAAGCAGCGTGGTA	522
humanTAS1R1	GCCGCCTGCTGAGCCCTTTCCTGGTGCCCATGATTAGCTATGCGGCCAGCAGCAGCAGACG	
mouseTaslr3	GGCAAGTTCTTCAGCTTCTTCCTCATGCCACAGGTCAGCTATAGTGCCAGCATGGATCGG	519
ratTas1r3	GGCAAGTTCTTCAGCTTCTTCCTCATGCCACAGGTCAGCTATAGTGCCAGCATGGATCGG GGCAAGTTCTTCAGCTTCTTCCTCATGCCACAGGTCAGCTATAGTGCCAGCATGGATCGG	519
		-
catTas1r3	GGCAAGTTCTTCAGCTTCCTTGTGCCTCAGGTCAGCTACGGCGCCAGCACCGACCG	531
humanTAS1R3	GGCAAGTTCTTCAGCTTCTTCCTCATGCCCCAGGTCAGCTACGGTGCTAGCATGGAGCTG  * * * * * * * * * * * * * * * * * * *	519
	4	
mouseTas1r2	CTGCGAGACAAGCGGCGCTTCCCTGCCATGCTGCGCACTGTGCCCAGCGCCACCCAC	579
ratTaslr2	CTGCGGGACAAGCGGCACTTCCCTAGCATGCTACGCACAGTGCCCAGCGCCACCCAC	579
humanTAS1R2	CTGCGAGACAAGGTGCGCTTCCCGGCTTTGCTGCGTACCACACCCAGCGCCGACCACCAC	570
mouseTas1r1	CTCAGTGGGAAGCGCAAGTTCCCGTCCTTCTTGCGCACCATCCCCAGCGATAAGTACCAG	588
ratTas1r1	CTCAGTGCCAAGCGCAAGTTCCCGTCTTTCCTTCGTACCGTCCCCAGTGACCGGCACCAG	582
humanTAS1R1	CTCAGCGTGAAGCGGCAGTATCCCTCTTTCCTGCGCACCATCCCCAATGACAAGTACCAG	585
mouseTas1r3	CTAAGTGACCGGGAAACGTTTCCATCCTTCTTCCGCACAGTGCCCAGTGACCGGGTGCAG	579
ratTas1r3	CTAAGTGACCGGGAAACATTTCCATCCTTCTTCCGCACAGTGCCCAGTGACCGGGTGCAG	
catTas1r3	CTGAGCAACCGGGAGATCTTCCCGTCCTTCTTCCGCACGGTGCCCAGCGACCAGGTGCAG	
humanTAS1R3	CTGAGCGCCCGGGAGACCTTCCCCCTCCTTCTTCCGCACCGTGCCCAGCGACCGTGTGCAG	
	** * * * * * * * * * * * * * * * * * * *	0.5
mouseTas1r2	ATCGAGGCCATGGTGCAACTGATGGTTCACTTCCAGTGGAACTGGATCGTGGTGCTGGTG	
ratTas1r2	ATCGAGGCCATGGTGCAGCTGATGGTTCACTTCCAATGGAACTGGATTGTGGTGCTGGTG	
humanTAS1R2	GTCGAGGCCATGGTGCAGCTGATGCTGCACTTCCGCTGGAACTGGATCATTGTGCTGGTG	630
mouseTas1r1	GTGGAAGTCATAGTGCGGCTGCTGCAGAGCTTCGGCTGGGTCTGGATCTCGCTCG	648
ratTas1r1	GTGGAGGTCATGGTGCAGCTGCAGAGTTTTGGGTGGGTGTGGATCTCGCTCATTGGC	642
humanTAS1R1	GTGGAGACCATGGTGCTGCTGCAGAAGTTCGGGTGGACCTGGATCTCTCTGGTTGGC	645
mouseTas1r3	CTGCAGGCAGTTGTGACTCTGTTGCAGAACTTCAGCTGGAACTGGGTGGCCGCCTTAGGG	
ratTas1r3	CTGCAGGCCGTTGTGACACTGTTGCAGAATTTCAGCTGGAACTGGGTGGCTGCCTTAGGT	639
catTaslr3	GTGGCGGCCATGGTGGAGCTGCTGGAGGAGCTCGGCTGGAACTGGGTGGCGGCGGTGGGT	
humanTAS1R3	CTGACGGCCGCGGAGCTGCTGCAGGAGTTCGGCTGGAACTGGGTGGCCGCCCTGGGC	
	* * *** ** * *** *** * * * * * * * * * *	

Docket No.: MON-3042

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Figur 1C

mouseTas1r2	AGCGATGACGATTATGGCCGAGAGAACAGCCACCTGCTGAGCCAGCGTCTGACCAACACT	699
ratTas1r2	AGCGACGATTACGGCCGCGAGAACAGCCACCTGTTGAGCCAGCGTCTGACCAAAACG	
humanTAS1R2	AGCAGCGACACCTATGGCCGCGACAATGGCCAGCTGCTTGGCGAGCGCGCGC	
	AGCTATGGTGACTACGGGCAGCTGGGCGTACAGGGCGCTGGAGGAGCACCTCCA	
mouseTaslr1		
ratTasir1	AGCTACGGTGATTACGGGCAGCTGGGTGTGCAGGCGCTGGAGGAGCTGGCCGTGCCC	
humanTAS1R1	AGCAGTGACGACTATGGGCAGCTAGGGGTGCAGGCACTGGAGAACCAGGCCACTGGT	
mouseTas1r3	AGTGATGATGACTATGGCCGGGAAGGTCTGAGCATCTTTTCTAGTCTGGCCAATGCA	
ratTas1r3	AGTGATGATGACTATGGCCGGGAAGGTCTGAGCATCTTTTCTGGTCTGGCCAACTCA	696
catTas1r3	AGTGACGACGAGTATGGCCGGCAGGGCCTGAGCCTCTTCTCCGGCCTGGCCAGCGCC	708
humanTAS1R3	AGCGACGACGAGTACGGCCGGCAGGGCCTGAGCATCTTCTCGGCCCTGGCCGCGGCA	696
	** * ** ** * * * * * *	
mouseTas1r2	GGCGATATCTGCATTGCCTTCCAGGAGGTTCTGCCTGTACCAGAACCCAACCAGGCCGTG	759
ratTas1r2	AGCGACATCTGCATTGCCTTCCAGGAGGTTCTGCCCATACCTGAGTCCAGCCAG	
humanTAS1R2	CGCGACATCTGCATCGCCTTCCAGGAGACGCTGCCCACACTGCAGCCCAACCAGAACATG	
mouseTasir1	CGGGGCATCTGCGTCGCCTTCAAGGACGTGGTGCCTCTCTCCGCCCAGGCGGG	
ratTas1r1	CGGGGCATCTGCGTCGCCTTCAAGGACATCGTGCCTTTCTCTGCCCGGGTGGG	
humanTAS1R1	CAGGGGATCTGCATTGCTTTCAAGGACATCATGCCCTTCTCTGCCCAGGTGGG	
mouseTaslr3	CGAGGTATCTGCATCGCACATGAGGGCCTGGTGCCACAACATGACACTAGTGGC	
ratTas1r3	CGAGGTATCTGCATTGCACACGAGGGCCTGGTGCCACAACATGACACTAGTGGC	
catTas1r3	AGGGCATCTGCATCGCGCATGAGGGCCTGGTGCCACTGC-CGCCAGGCAGC	
humanTAS1R3	CGCGGCATCTGCGCACGAGGGCCTGGTGCCGCTGCCCCGTGCCGATGAC	750
	* ***** * ** ***	
mouseTas1r2	AGGCCTGAGGAGCAGGACCAACTGGACAACATCCTGGACAAGCTGCGGCGGACCTCGGCG	819
ratTas1r2	AGGTCCGAGGAGCAGAGACAACTGGACAACATCCTGGACAAGCTGCGGCGGACCTCGGCG	
humanTAS1R2	ACGTCAGAGGGGCGCCAGCGCTGGTGACCATTGTGGACAAGCTGCAGCAGAGCACAGCG	
mouseTas1r1	TGACCCAAGGATGCAGCGCATGATGCTGCGTCTGGCTCGAGCCAGGACC	
ratTaslr1		
	TGACCCGAGGATGCAGAGCATGATGCAGCATCTGGCTCAGGCCAGGACC	
humanTAS1R1	CGATGAGAGGATGCAGTGCCTCATGCGCCACCTGGCCCAGGCCGGGGCC	
mouseTas1r3	CAACAGTTGGGCAAGGTGCTGGATGTACTACGCCAAGTGAACCAAAGTAAAGT	
ratTas1r3	CAACAATTGGGCAAGGTGGTGGATGTGCTACGCCAAGTGAACCAAAGCAAAG	
catTas1r3	CTGCGGCTGGGCGCCCTACAGGGCCTGCTGCGCCAGGTGAACCAGAGCAGCGTG	
humanTAS1R3	TCGCGGCTGGGGAAGGTGCAGGACGTCCTGCACCAGGTGAACCAGAGCAGCGTG	804
	**	
mouseTas1r2	CGTGTGGTGGTGATATTCTCGCCAGAGCTGAGCCTGCACAACTTCTTCCGCGAGGTGC	877
ratTas1r2	CGCGTCGTGGTGGTGTTCTCGCCCGAGCTGAGCCTGTATAGCTTCTTTCACGAGGTGC	
humanTAG1D2	CCCCTCCTCCTCCTCCTCCTCCTCCTCCTCCTCCTCCTC	
humanTAS1R2	CGCGTCGTGGTCGTGTTCTCGCCCGACCTGACCCTGTACCACTTCTTCAATGAGGTGC	865
mouseTas1r1	ACCGTGGTCGTGGTCTTCTCTAACCGGCACCTGGCTGGAGTGTTCTTCAGGTCTGTGG	865 865
mouseTaslr1 ratTaslr1	ACCGTGGTCGTGGTCTTCTCTAACCGGCACCTGGCTGGAGTGTTCTTCAGGTCTGTGG ACCGTGGTTGTGGTCTTCTCTAACCGGCACCTGGCTAGAGTGTTCTTCAGGTCCGTGG	865 865 859
mouseTaslrl ratTaslrl humanTASlRl	ACCGTGGTCGTGGTCTTCTCTAACCGGCACCTGGCTGGAGTGTTCTTCAGGTCTGTGG ACCGTGGTTGTGGTCTTCTCTAACCGGCACCTGGCTAGAGTGTTCTTCAGGTCCGTGG ACCGTCGTGGTTGTTTTTTCCAGCCGGCAGTTGGCCAGGGTGTTTTTCGAGTCCGTGG	865 865 859 862
mouseTaslr1 ratTaslr1 humanTASlR1 mouseTaslr3	ACCGTGGTCGTGGTCTTCTCTAACCGGCACCTGGCTGGAGTGTTCTTCAGGTCTGTGG ACCGTGGTTGTGGTCTTCTCTAACCGGCACCTGGCTAGAGTGTTCTTCAGGTCCGTGG ACCGTCGTGGTTGTTTTTTCCAGCCGGCAGTTGGCCAGGGTGTTTTTCGAGTCCGTGG CAAGTGGTGGTGCTGTTTGCCTCTGCCCGTGCTGTCTACTCCCTTTTTAGTTACAGCA	865 865 859 862 862
mouseTaslr1 ratTaslr1 humanTASlR1 mouseTaslr3 ratTaslr3	ACCGTGGTCGTGGTCTTCTCTAACCGGCACCTGGCTGGAGTGTTCTTCAGGTCTGTGG ACCGTGGTTGTGTCTCTCTAACCGGCACCTGGCTAGAGTGTTCTTCAGGTCCGTGG ACCGTCGTGGTTGTTTTTCCAGCCGGCAGTTGGCCAGGGTGTTTTTCGAGTCCGTGG CAAGTGGTGGTGCTGTTTGCCTCTGCCCGTGCTGTCTACTCCCTTTTTAGTTACAGCA CAGGTGGTGGTGCTGTTTGCATCTGCCCGTGCTGTCTACTCCCTTTTTAGCTACAGCA	865 865 859 862 862 862
mouseTaslr1 ratTaslr1 humanTASlR1 mouseTaslr3 ratTaslr3 catTaslr3	ACCGTGGTCGTGGTCTTCTCTAACCGGCACCTGGCTGGAGTGTTCTTCAGGTCTGTGG ACCGTGGTTGTGTCTCTTAACCGGCACCTGGCTAGAGTGTTCTTCAGGTCCGTGG ACCGTCGTGGTTGTTTTTCCAGCCGGCAGTTGGCCAGGGTGTTTTTCGAGTCCGTGG CAAGTGGTGGTGCTGTTTGCCTCTGCCCGTGCTGTCTACTCCCTTTTTAGTTACAGCA CAGGTGGTGGTGCTGTTTTGCATCTGCCCGTGCTGTCTACTCCCTTTTTAGCTACAGCA CAGGTGGTGGTGCTGTTCTCCTCCGCCCACGCGGCCCCACCCTCTTCAGCTACAGCA	865 865 859 862 862 862 871
mouseTaslr1 ratTaslr1 humanTASlR1 mouseTaslr3 ratTaslr3	ACCGTGGTCGTGGTCTTCTCTAACCGGCACCTGGCTGGAGTGTTCTTCAGGTCTGTGG ACCGTGGTTGTGGTCTTCTCTAACCGGCACCTGGCTAGAGTGTTCTTCAGGTCCGTGG ACCGTCGTGGTTGTTTTTCCAGCCGGCAGTTGGCCAGGGTGTTTTTCGAGTCCGTGG CAAGTGGTGGTGCTGTTTGCCTCTGCCCGTGCTGTCTACTCCCTTTTTAGTTACAGCA CAGGTGGTGGTGCTGTTTCCCTCCGCCACGCGGCCCGCACCCTCTTCAGCTACAGCA CAGGTGGTGGTGCTGTTCCCTCCGTGCACGCCCCCACCCCCTCTTCAACTACAGCA	865 865 859 862 862 862 871
mouseTaslr1 ratTaslr1 humanTASlR1 mouseTaslr3 ratTaslr3 catTaslr3	ACCGTGGTCGTGGTCTTCTCTAACCGGCACCTGGCTGGAGTGTTCTTCAGGTCTGTGG ACCGTGGTTGTGTCTCTTAACCGGCACCTGGCTAGAGTGTTCTTCAGGTCCGTGG ACCGTCGTGGTTGTTTTTCCAGCCGGCAGTTGGCCAGGGTGTTTTTCGAGTCCGTGG CAAGTGGTGGTGCTGTTTGCCTCTGCCCGTGCTGTCTACTCCCTTTTTAGTTACAGCA CAGGTGGTGGTGCTGTTTTGCATCTGCCCGTGCTGTCTACTCCCTTTTTAGCTACAGCA CAGGTGGTGGTGCTGTTCTCCTCCGCCCACGCGGCCCCACCCTCTTCAGCTACAGCA	865 865 859 862 862 862 871
mouseTaslr1 ratTaslr1 humanTASlR1 mouseTaslr3 ratTaslr3 catTaslr3	ACCGTGGTCGTGGTCTTCTCTAACCGGCACCTGGCTGGAGTGTTCTTCAGGTCTGTGG ACCGTGGTTGTGGTCTTCTCTAACCGGCACCTGGCTAGAGTGTTCTTCAGGTCCGTGG ACCGTCGTGGTTGTTTTTCCAGCCGGCAGTTGGCCAGGGTGTTTTTCGAGTCCGTGG CAAGTGGTGGTGCTGTTTGCCTCTGCCCGTGCTGTCTACTCCCTTTTTAGTTACAGCA CAGGTGGTGGTGCTGTTTGCATCTGCCCGTGCTGTCTACTCCCTTTTTAGCTACAGCA CAGGTGGTGGTGCTGTTCTCCTCCGCCCACGCGCCCCCACCCTCTTCAGCTACAGCA CAGGTGGTGCTGCTTCTCCTCCGTGCACCCCCCCCCCC	865 865 859 862 862 862 871 862
mouseTaslr1 ratTaslr1 humanTAslR1 mouseTaslr3 ratTaslr3 catTaslr3 humanTASlR3	ACCGTGGTCGTGGTCTTCTCTAACCGGCACCTGGCTGGAGTGTTCTTCAGGTCTGTGGACCGTGGTTGTGTGTTTTTTTTCTAGTCTTCTTTTTTTT	865 865 859 862 862 862 871 862
mouseTaslr1 ratTaslr1 humanTASlR1 mouseTaslr3 ratTaslr3 catTaslr3 humanTASlR3 mouseTaslr2 ratTaslr2	ACCGTGGTCGTGGTCTTCTCTAACCGGCACCTGGCTGAGTG-TTCTTCAGGTCTGTGG ACCGTGGTTGTGGTCTTCTCTAACCGGCACCTGGCTAGAGTG-TTCTTCAGGTCCGTGG ACCGTCGTTGTTTTTTCCAGCCGGCACTTGGCCAGGGTG-TTTTTTCGAGTCCGTGG CAGTGGTGGTGCTGTTTGCCTCTGCCCGTGCTGTCTCTCTC	865 865 859 862 862 862 871 862 937 937
mouseTaslr1 ratTaslr1 humanTASlR1 mouseTaslr3 ratTaslr3 catTaslr3 humanTASlR3  mouseTaslr2 ratTaslr2 humanTASlR2	ACCGTGGTCGTGGTCTTCTCTAACCGGCACCTGGCTGAGTG-TTCTTCAGGTCTGTGG ACCGTGGTTGTGGTCTTCTCTAACCGGCACCTGGCTAGAGTG-TTCTTCAGGTCCGTGG ACCGTCGTTGTTTTTTCCAGCCGGCAGTTGGCCAGGGTG-TTTTTTCGAGTCCGTGG CAGTGGTGGTGCTGTTTTGCCTCTGCCCGTGCTGTCTACTCCCTTTTTAGCTACAGCA CAGGTGGTGGTGTTTCCCTCCGCCACGCGGCCCACCCTCTTCAGCTACAGCA CAGGTGGTGCTGTTCCCTCCGTGCACGCCCCCCCCCCTCTTCAACTACAGCA ** ** * * * * * * * * * * * * * * * *	865 865 859 862 862 862 871 862 937 937 925
mouseTaslr1 ratTaslr1 humanTASlR1 mouseTaslr3 ratTaslr3 catTaslr3 humanTASlR3  mouseTaslr2 ratTaslr2 humanTASlR2 mouseTaslr1	ACCGTGGTCGTGGTCTTCTCTAACCGGCACCTGGCTGGAGTGTTCTTCAGGTCTGTGGACCGTGGTTGTGTGTCTTCTCTAACCGGCACCTGGCTAGAGTGTTCTTCAGGTCTGTGGACCGTGGTTGTTTTTTCTTCTTCTTCTTCTTCTTCTTCTTC	865 865 869 862 862 871 862 937 937 925 925
mouseTaslr1 ratTaslr1 humanTASlR1 mouseTaslr3 ratTaslr3 catTaslr3 humanTASlR3  mouseTaslr2 ratTaslr2 humanTASlR2 mouseTaslr1 ratTaslr1	ACCGTGGTCGTGGTCTTCTCTAACCGGCACCTGGCTGGAGTG-TTCTTCAGGTCTGTGGACCGTGGTTGTGTGTCTTCTCTAACCGGCACCTGGCTAGAGTG-TTCTTCAGGTCTGTGGACCGTGGTTGTTTTTTTTTT	865 865 869 862 862 871 862 937 937 925 925 919
mouseTaslr1 ratTaslr1 humanTAslR1 mouseTaslr3 ratTaslr3 catTaslr3 humanTASlR3  mouseTaslr2 ratTaslr2 humanTASlR2 mouseTaslr1 ratTaslr1 humanTASlR1	ACCGTGGTCGTGGTCTTCTCTAACCGGCACCTGGCTGAGTG-TTCTTCAGGTCTGTGGACCGGTGGTTGTGTGTTTTTTTTCTTCTTCTTTTTTTT	865 865 869 862 862 862 871 862 937 937 925 925 919
mouseTaslr1 ratTaslr1 humanTASlR1 mouseTaslr3 ratTaslr3 catTaslr3 humanTASlR3  mouseTaslr2 ratTaslr2 humanTASlR2 mouseTaslr1 ratTaslr1 humanTASlR1 mouseTaslr1 mouseTaslr1	ACCGTGGTCGTGGTCTTCTCTAACCGGCACCTGGCTGAGTG-TTCTTCAGGTCTGTGG ACCGTGGTTGTGGTCTTCTCTAACCGGCACCTGGCTAGAGTG-TTCTTCAGGTCCGTGG ACCGTCGTTGTTTTTTCCAGCCGGCAGTTGGCCAGGGTG-TTCTTCAGGTCCGTGG ACCGTCGTGGTTGTTTTTCC-CTCCGCCGTGCTGTCTCTCTC	865 865 869 862 862 862 871 862 937 937 925 925 919 922 922
mouseTaslr1 ratTaslr1 humanTASlR1 mouseTaslr3 ratTaslr3 catTaslr3 humanTASlR3  mouseTaslr2 ratTaslr2 humanTASlR2 mouseTaslr1 ratTaslr1 humanTASlR1 mouseTaslr1 ratTaslr3 ratTaslr3	ACCGTGGTCGTGGTCTTCTCTAACCGGCACCTGGCTGAGTG-TTCTTCAGGTCTGTGGACCGTGGTTGTGTGTCTTCTTCAGGTCTTTCTT	865 865 869 862 862 862 871 862 937 937 925 925 919 922 922
mouseTaslr1 ratTaslr1 humanTASlR1 mouseTaslr3 ratTaslr3 catTaslr3 humanTASlR3  mouseTaslr2 ratTaslr2 humanTASlR2 mouseTaslr1 ratTaslr1 humanTASlR1 mouseTaslr1 ratTaslr1 humanTASlR3 catTaslr3 catTaslr3	ACCGTGGTCGTGGTCTTCTCTAACCGGCACCTGGCTGAGTG-TTCTTCAGGTCTGTGGACCGTGGTTGTGTGTCTTCTTCAACCGGCACCTGGCTAGAGTG-TTCTTCAGGTCTGTGGACCGTGGTTGTTTTTTCTTCTTCTTCTTCTTCTTCTTCTTC	865 865 865 862 862 862 861 862 937 937 925 925 919 922 922 931
mouseTaslr1 ratTaslr1 humanTASlR1 mouseTaslr3 ratTaslr3 catTaslr3 humanTASlR3  mouseTaslr2 ratTaslr2 humanTASlR2 mouseTaslr1 ratTaslr1 humanTASlR1 mouseTaslr1 ratTaslr3 ratTaslr3	ACCGTGGTCGTGGTCTTCTCTAACCGGCACCTGGCTGAGTG-TTCTTCAGGTCTGTGGACCGTGGTTGTGTGTCTTCTTCTAACCGGCACCTGGCTAGAGTG-TTCTTCAGGTCTGTGGACCGTGGTTGTTTTTTTTTT	865 865 865 862 862 862 861 862 937 937 925 925 919 922 922 931
mouseTaslr1 ratTaslr1 humanTASlR1 mouseTaslr3 ratTaslr3 catTaslr3 humanTASlR3  mouseTaslr2 ratTaslr2 humanTASlR2 mouseTaslr1 ratTaslr1 humanTASlR1 mouseTaslr1 ratTaslr1 humanTASlR3 catTaslr3 catTaslr3	ACCGTGGTCGTGGTCTTCTCTAACCGGCACCTGGCTGAGTG-TTCTTCAGGTCTGTGGACCGTGGTTGTGTGTCTTCTTCAACCGGCACCTGGCTAGAGTG-TTCTTCAGGTCTGTGGACCGTGGTTGTTTTTTCTTCTTCTTCTTCTTCTTCTTCTTC	865 865 865 862 862 862 861 862 937 937 925 925 919 922 922 931
mouseTaslr1 ratTaslr1 humanTASlR1 mouseTaslr3 ratTaslr3 catTaslr3 humanTASlR3  mouseTaslr2 ratTaslr2 humanTASlR2 mouseTaslr1 ratTaslr1 humanTASlR1 mouseTaslr1 ratTaslr1 humanTASlR1 mouseTaslr3 ratTaslr3 ratTaslr3 humanTASlR3	ACCGTGGTCGTGGTCTTCTCTAACCGGCACCTGGCTGAGTG-TTCTTCAGGTCTGTGGACCGTGGTTGTGTGTCTTCTTCTAACCGGCACCTGGCTAGAGTG-TTCTTCAGGTCTGTGGACCGTGGTTGTTTTTTCTTCTTCTTCTTCTTCTTCTTCTTC	865 865 865 862 862 862 871 862 937 937 925 925 919 922 922 922 922 931
mouseTaslr1 ratTaslr1 humanTASlR1 mouseTaslr3 ratTaslr3 catTaslr3 humanTASlR3  mouseTaslr2 ratTaslr2 humanTASlR2 mouseTaslr1 ratTaslr1 humanTASlR1 mouseTaslr3 ratTaslr3 catTaslr3 catTaslr3 humanTASlR3	ACCGTGGTCGTGGTCTTCTCTAACCGGCACCTGGCTGAGTG-TTCTTCAGGTCTGTGGACCGTGGTTGTGTGTCTTCTTCAGGTCTTTTCTCAGGTCTTTGGACCGTGGTTGGT	865 865 865 862 862 862 871 862 937 925 925 929 922 922 931 922
mouseTaslr1 ratTaslr1 humanTASlR1 mouseTaslr3 ratTaslr3 catTaslr3 humanTASlR3  mouseTaslr2 ratTaslr2 humanTASlR2 mouseTaslr1 ratTaslr1 humanTASlR1 mouseTaslr3 ratTaslr3 catTaslr3 catTaslr3 humanTASlR3  mouseTaslr3 ratTaslr3 catTaslr3 catTaslr3 humanTASlR3	ACCGTGGTCGTGGTCTTCTCTAACCGGCACCTGGCTGAGTG-TTCTTCAGGTCTGTGGACCGTGGTTGTGTGTCTTCTTCAGGTCTTTTTCAGGTCTTTGTGAGCCGTGGTTGGT	865 865 865 862 862 862 871 862 937 925 925 929 922 931 922 931 922
mouseTaslr1 ratTaslr1 humanTAslR1 mouseTaslr3 ratTaslr3 catTaslr3 humanTAslR3  mouseTaslr2 ratTaslr2 humanTAslR2 mouseTaslr1 ratTaslr1 humanTAslR1 mouseTaslr3 ratTaslr3 ratTaslr3 ratTaslr3 ratTaslr3 ratTaslr3 humanTAslR3	ACCGTGGTCGTGGTCTTCTCTAACCGGCACCTGGCTGAGTG-TTCTTCAGGTCTGTGGACCGTGGTTGTGTGTGTTTTTTTT	865 865 862 862 862 871 862 937 937 925 919 922 922 922 922 922 931 922
mouseTaslr1 ratTaslr1 humanTASlR1 mouseTaslr3 ratTaslr3 catTaslr3 humanTASlR3  mouseTaslr2 ratTaslr2 humanTASlR2 mouseTaslr1 ratTaslr1 humanTASlR1 mouseTaslr3 ratTaslr3 ratTaslr3 catTaslr3 ratTaslr3 catTaslr3 ratTaslr3 ratTaslr3 catTaslr3 humanTASlR3	ACCGTGGTCGTGGTCTTCTCTAACCGGCACCTGGCTGAGTG-TTCTTCAGGTCTGTGGACCGTGGTTGTGTGTCTTCTTCTAACCGGCACCTGGCTAGAGTG-TTCTTCAGGTCGTGGACCGTGGTTGTTTTTTCTTCTTCTTCTTCTTCTTCTTCTTC	865 865 862 862 862 871 862 937 925 925 919 922 922 931 922 997 997 9985 985
mouseTaslr1 ratTaslr1 humanTASlR1 mouseTaslr3 ratTaslr3 catTaslr3 humanTASlR3  mouseTaslr2 ratTaslr2 humanTASlR2 mouseTaslr1 ratTaslr1 humanTASlR1 mouseTaslr3 ratTaslr3 catTaslr3 catTaslr3 catTaslr3 humanTASlR3  mouseTaslr3 ratTaslr3 ratTaslr3 catTaslr3 humanTASlR3	ACCGTGGTCGTGGTCTTCTCTAACCGGCACCTGGCTGAGTG-TTCTTCAGGTCTGTGACCGTGGTTGTGTGTCTTCTTCTAACCGGCACCTGGCTAGAGTG-TTCTTCAGGTCGTGGACCGTGGTTGTTTTTTCTTCTTCTTCTTCTTCTTCTTCTTTTTT	865 865 862 862 862 871 862 937 937 925 925 922 922 931 922 937 985 979
mouseTaslr1 ratTaslr1 humanTASlR1 mouseTaslr3 ratTaslr3 catTaslr3 humanTASlR3  mouseTaslr2 ratTaslr2 humanTASlR2 mouseTaslr1 ratTaslr1 humanTASlR1 mouseTaslr3 ratTaslr3 catTaslr3 catTaslr3 catTaslr3 humanTASlR3  mouseTaslr2 ratTaslr2 humanTASlR3	ACCGTGGTCGTGGTCTTCTCTAACCGGCACCTGGCTGAGTG-TTCTTCAGGTCTGTGGACCGTGGTTGTGTGTCTTCTTAACCGGCACCTGGCTAGAGTG-TTTCTTCAGGTCTGTGGACCGTGGTTTTTTTTTT	865 865 862 862 862 871 862 937 937 925 922 922 931 922 931 922 937 985 985
mouseTaslr1 ratTaslr1 humanTASlR1 mouseTaslr3 ratTaslr3 catTaslr3 humanTASlR3  mouseTaslr2 ratTaslr2 humanTASlR2 mouseTaslr1 ratTaslr1 humanTASlR1 mouseTaslr3 ratTaslr3 ratTaslr3 ratTaslr3 catTaslr3 humanTASlR3  mouseTaslr2 ratTaslr2 humanTASlR2 mouseTaslr2 ratTaslr2 humanTASlR2 mouseTaslr1 ratTaslr1 humanTASlR1 mouseTaslr3 mouseTaslr3 mouseTaslr3 humanTASlR2 mouseTaslr1 ratTaslr1 humanTASlR1 mouseTaslr3	ACCGTGGTCGTGGTCTTCTCTAACCGGCACCTGGCTGAGTG-TTCTTCAGGTCTGTGACCGTGGTTGTGTGTGTTTTTTTTT	865 865 862 862 862 871 937 925 925 919 922 931 922 931 922 937 985 979 985 979 982 982
mouseTaslr1 ratTaslr1 humanTAslR1 mouseTaslr3 ratTaslr3 catTaslr3 humanTAslR3  mouseTaslr2 ratTaslr2 humanTAslR1 mouseTaslr1 ratTaslr1 humanTAslR1 mouseTaslr3 ratTaslr3 ratTaslr3 catTaslr3 catTaslr3 humanTAslR3  mouseTaslr2 ratTaslr2 humanTAslR2 mouseTaslr2 ratTaslr3 ratTaslr3 ratTaslr3 ratTaslr3 ratTaslr3 ratTaslr3 ratTaslr3	ACCGTGGTCGTGGTCTTCTCTAACCGGCACCTGGCTGAGTG-TTCTTCAGGTCTGTGGACCGTGGTTGTGTGTCTTCTTAACCGGCACCTGGTTAGAGTG-TTCTTCAGGTCGTGGACCGTGGTTGTTTTTTTTTT	865 865 862 862 862 871 862 937 925 925 919 922 922 931 922 997 985 979 982 982 982
mouseTaslr1 ratTaslr1 humanTASlR1 mouseTaslr3 ratTaslr3 catTaslr3 humanTASlR3  mouseTaslr2 ratTaslr2 humanTASlR2 mouseTaslr1 ratTaslr1 humanTASlR1 mouseTaslr3 ratTaslr3 catTaslr3 catTaslr3 humanTASlR3  mouseTaslr1 ratTaslr1 humanTASlR3  mouseTaslr2 ratTaslr2 humanTASlR1 mouseTaslr2 ratTaslr3 ratTaslr3 catTaslr3 ratTaslr1 humanTASlR1 mouseTaslr1 ratTaslr1 humanTASlR1 mouseTaslr3 ratTaslr3 catTaslr3 catTaslr3	ACCGTGGTCGTGGTCTTCTCTAACCGGCACCTGGCTGAGTG-TTCTTCAGGTCTGTGACCGTGGTTGTGTGTCTTCTTCTAACCGGCACCTGGCTAGAGTG-TTTCTTCAGGTCGTGGACCGTGGTTTGTTTTTTCTTCTTTTTTTT	865 865 862 862 862 871 862 937 937 925 925 922 922 931 922 937 985 979 982 982 997
mouseTaslr1 ratTaslr1 humanTAslR1 mouseTaslr3 ratTaslr3 catTaslr3 humanTAslR3  mouseTaslr2 ratTaslr2 humanTAslR1 mouseTaslr1 ratTaslr1 humanTAslR1 mouseTaslr3 ratTaslr3 ratTaslr3 catTaslr3 catTaslr3 humanTAslR3  mouseTaslr2 ratTaslr2 humanTAslR2 mouseTaslr2 ratTaslr3 ratTaslr3 ratTaslr3 ratTaslr3 ratTaslr3 ratTaslr3 ratTaslr3	ACCGTGGTCGTGGTCTTCTCTAACCGGCACCTGGCTGAGTG-TTCTTCAGGTCTGTGGACCGTGGTTGTGTGTCTTCTTAACCGGCACCTGGTTAGAGTG-TTCTTCAGGTCGTGGACCGTGGTTGTTTTTTTTTT	865 865 862 862 862 871 862 937 937 925 925 922 922 931 922 937 985 979 982 982 997

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Title: TASTE RECEPTOR OF THE TIR FAMILY FROM DOMESTIC CAT Inventors: Xia Ll, Weihua Ll, Danielle R. REED, Alexander Alexeyevich BACHMANOV, Joseph G. BRAND

Attorney: Felicity E. Groth Sheet 4 of 13

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Figure 1D

humanTAS1R1 mouseTas1r3

humanTAS1R3

ratTas1r3 catTas1r3

mouseTas1r2 GGGTGTCCATCCCTGGCTTCAGCCAGTTCCGAGTGCGCCAC---GACAAGCCAG----- 1048 ratTas1r2 GGGTGTCCATCCCTGGCTTCAGTCAGTTCCGAGTGCGCCGT---GACAAGCCAG----- 1048 humanTAS1R2 GCGTGCCCATCCCGGGCTTCAGTGAGTTCCGCGAGTGGGGC---CCACAGGCTG----- 1036 mouseTas1r1 AGAGACAAGTCCCTGGCCTGAAGGAGTTTGAAGAGTCCTAT---GTCCAGGCAGTGATGG 1042 AGAGACAAGTCCCTGGGCTGAAGGAGTTTGAGGAGTCTTAT---GTCAGGGCTGTAACAG 1036 ratTas1r1 AGAGGGCTGTCCCTGGCCTGAAGGCGTTTGAAGAAGCCTAT---GCCCGGGCAGACAAGA 1039 humanTAS1R1 GTGCCCTACTGCCTGAATTTTCCCATTATGTGGAGACTCACCTTGCCCTGGCCGCTGACC 1042 mouseTas1r3 ratTas1r3 GTGCCCTACTGCCTGAATTTTCCCATTATGTGGAGACTCGCCTTGCCCTAGCTGCTGACC 1042 catTas1r3 GCGCCCGATGCCGGAGTTCCCATCCTACGTGCGGACCCGCCTGGCCCTGGCCGCTGACC 1051 humanTAS1R3 GTGCCCAGCTGCACGAGTTCCCCCAGTACGTGAAGACGCACCTGGCCCTGGCCACCGACC 1042 mouseTas1r2 -AGTATCCCATGCCTAACGAGACCAGCC----TGAGGACTACCTGTAACCAGGAC 1098 -GGTATCCCGTGCCTAACACGACCAACC-----TGCGGACGACCTGCAACCAGGAC 1098 ratTas1r2 -GGCCGCCACCCTCAGCAGGACCAGCC-----AGAGCTATACCTGCAACCAGGAG 1086 humanTAS1R2 GTGCTCCCAGAACTTGCCCAGAGGGGTC-----C-TGGTGCGGCACTAACCAGCTG 1092 mouseTas1r1 CTGCTCCCAGCGCTTGCCCGGAGGGGTC-----C-TGGTGCAGCACTAACCAGCTG 1086 ratTas1r1 AGGCCCCTAGGCCTTGCCACAAGGGCTC-----C-TGGTGCAGCAGCAATCAGCTC 1089 humanTAS1R1 CAGCATTCTGTGCCTCACTGAATGCGGA~--GTTGGATCTGGAGGAACATGTGATGGGGC 1099 mouseTas1r3 CAACATTCTGTGCCTCCCTGAAAGCTGA---GTTGGATCTGGAGGAGCGCGTGATGGGGC 1099 ratTaslr3 CTGCCTTCTGCGCCTCGCTGGACGCTGAACAGCCAGGCCTGGAGGAGCACGTGGTGGGGC 1111 catTaslr3 humanTAS1R3 CGGCCTTCTGCTCTGCCCTGGGCGAGAGGGAGGAGGACGTGGTGGGCC 1102 TGTGACGCCTGCATGAACATCACCGAGTCCTTTAACAACGTTC----- 1141 mouseTas1r2 TGTGACGCCTGCTTGAACACCACCAAGTCCTTCAACAACATCC------ 1141 ratTas1r2 TGCGACAACTGCCTGAACGCCACCTTGTCCTTCAACACCATTC----- 1129 humanTAS1R2 TGCAGGGAGTGTCACGCTTTCACGACATGGAACATGCCCGAGC----- 1135 mouseTas1r1 TGCCGGGAGTGCCACACGTTCACGACTCGTAACATGCCCACGC----- 1129 ratTas1r1 TGCAGAGAATGCCAAGCTTTCATGGCACACACGATGCCCAAGC----- 1132 humanTAS1R1 AACGCTGTCCACGGTGTGACGACATCATGCTGCAGAACCTATCATCTGGGCTGTTGCAGA 1159 mouseTas1r3 CACGCTGTTCACAATGTGACTACATCATGCTACAGAACCTGTCATCTGGGCTGATGCAGA 1159 ratTas1r3 catTas1r3 CACGCTGCCCCCAATGTGACCACGTCACGCTAGAGAACCT------ 1151 AGCGCTGCCCGCAGTGTGACTGCATCACGCTGCAGAACGT----- 1142 humanTAS1R3 -TCATGCTTTCGGGGGAGCGTGTGGTC----TACAGTGTGTACTCGGCCGTCTACGCGG 1195 mouseTas1r2 -TTATACTTTCGGGGGAGCGCGTGGTC----TACAGCGTGTACTCGGCAGTTTACGCGG 1195 ratTas1r2 -TCAGGCTCTCTGGGGAGCGTGTCGTC----TACAGCGTGTACTCTGCGGTCTATGCTG 1183 humanTAS1R2 -TTGGAGCCTTCTCCATGAGCGCTGCC----TACAATGTGTATGAGGCTGTGTATGCTG 1189 mouseTas1rl -TTGGAGCCTTCTCCATGAGTGCCGCC----TACAGAGTGTATGAGGCTGTGTACGCTG 1183 ratTas1r1 -TCAAAGCCTTCTCCATGAGTTCTGCC----TACAACGCATACCGGGCTGTGTATGCGG 1186 humanTAS1R1 ACCTATCAGCTGGGCAATTGCACCACCAAATATTTGCAACCTATGCAGCTGTGTACAGTG 1219 mouseTas1r3 ACCTATCAGCTGGGCAGTTGCACCACCAAATATTTGCAACCTATGCAGCTGTGTACAGTG 1219 ratTas1r3 ----ATCTGCGGGGCTGCTGCACCACACACACCTCGCTGCCTACGCGGCTGTGTATGGCG 1207 catTas1r3 ----GAGCGCAGGGCTAAATCACCACCAGACGTTCTCTGTCTACGCAGCTGTGTATAGCG 1198 humanTAS1R3 TAGCCCACACCCTCCACAGACTCCTCCACTGCAACCAGGTCCGCTGCACCA---AGCAAA 1252 mouseTas1r2 TGGCCCATGCCCTCCACAGACTCCTCGGCTGTAACCGGGTCCGCTGCACCA---AGCAAA 1252 ratTaslr2 humanTAS1R2 TGGCCCATGCCCTGCACAGCCTCCTCGGCTGTGACAAAAGCACCTGCACCA---AGAGGG 1240 TGGCCCACGGCCTCCACCAGCTCCTGGGATGTACCTCTGGGACCTGTGCCA---GAGGCC 1246 mouseTas1r1 ratTas1r1 TGGCCCACGGCCTCCACCAGCTCCTGGGATGTACTTCTGAGATCTGTTCCA---GAGGCC 1240 TGGCCCATGGCCTCCACCAGCTCCTGGGCTGTGCCTCTGGAGCTTGTTCCA---GGGGCC 1243 humanTAS1R1 TGGCTCAAGCCCTTCACAACACCCTACAGTGCAATGTCTCACATTGCCACGTATCAGAAC 1279 mouseTas1r3 TGGCTCAGGCCCTTCACAACACCCTGCAGTGCAATGTCTCACATTGCCACACATCAGAGC 1279 ratTas1r3 TGGCCCAAGCCCTTCACAACACACTGCGCTGCAATGCCTCGGGCTGCCCCAGGCGGGAGC 1267 catTas1r3 TGGCCCAGGCCCTGCACAACACTCTTCAGTGCAACGCCTCAGGCTGCCCCGCGCAGGACC 1258 humanTAS1R3 mouseTas1r2 TCGTCTATCCATGGCAGCTACTCAGGGAGATCTGGCATGTCAACTTCACGCTCCTGGGCA 1312 AGGTCTACCCGTGGCAGCTACTCAGGGAGATCTGGCACGTCAACTTCACGCTCCTGGGTA 1312 ratTas1r2 humanTAS1R2 TGGTCTACCCCTGGCAGCTGCTTGAGGAGGTCTGGAAGGTCAACTTCACTCTCCTGGACC 1300 CAGTCTACCCCTGGCAGCTTCTTCAGCAGATCTACAAGGTGAATTTCCTTCTACATAAGA 1306 mouseTas1r1 CAGTCTACCCCTGGCAGCTTCTTCAGCAGATCTACAAGGTGAATTTTCTTCTACATGAGA 1300 ratTas1r1

GAGTCTACCCCTGGCAGCTTTTGGAGCAGATCCACAAGGTGCATTTCCTTCTACACAAGG 1303

ATGTTCTACCCTGGCAGCTCCTGGAGAACATGTACAATATGAGTTTCCATGCTCGAGACT 1339 CTGTTCAACCCTGGCAGCTCCTGGAGAACATGTACAATATGAGTTTCCGTGCTCGAGACT 1339

CTGTGCGGCCCTGGCAGCTCCTAGAGAACATGTACAACGTGAGCTTCCGTGCTCGCGGCC 1327

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Title: TASTE RECEPTOR OF THE TIR FAMILY FROM DOMESTIC CAT

Inventors: Xia Ll, Weihua LI, Danielle R. REED, Alexander Alexeyevich BACHMANOV,

Joseph G. BRAND Attorney: Felicity E. Groth Sheet 5 of 13

Phone: (215) 568-3100

### Figure 1E

humanTAS1R1

mouseTas1r3

ratTas1r3

catTas1r3 humanTAS1R3

mouseTas1r2 ACCAGCTCTTCTTCGACGAACAAGGGGACATGCCGATGCTCCTGGACATCATCCAGTGGC 1372 ratTas1r2 ACCGGCTCTTCTTTGACCAACAAGGGGACATGCCGATGCTCTTGGACATCATCCAGTGGC 1372 humanTAS1R2 ACCAAATCTTCTTCGACCCGCAAGGGGACGTGGCTCTGCACTTGGAGATTGTCCAGTGGC 1360 mouseTas1r1 AGACTGTAGCATTCGATGACAAGGGGGACCCTCTAGGTTATTATGACATCATCGCCTGGG 1366 ratTaslrl ATACTGTGGCATTTGATGACAACGGGGACACTCTAGGTTACTACGACATCATCGCCTGGG 1360 humanTAS1R1 ACACTGTGGCGTTTAATGACAACAGAGATCCCCTCAGTAGCTATAACATAATTGCCTGGG 1363 mouseTaslr3 TGACACTACAGTTTGATGCTGAAGGGAATGTAGACATGGAATATGACCTGAAGATGTGGG 1399 ratTas1r3 TGACACTGCAGTTTGATGCCAAAGGGAGTGTAGACATGGAATATGACCTGAAGATGTGGG 1399 catTaslr3 TGGCACTGCAGTTCGACGCCAGCGGGAACGTGAACGTGGATTACGACCTGAAACTGTGGG 1387 humanTAS1R3 TGCCGCTGCGGTTCGACAGCAGCGGAAACGTGGACATGGACTACGACCTGAAGCTGTGGG 1378 mouseTas1r2 AATGGGGCCTGAGCCAGAACCCCTTCCAAAGCATCGCCTCCTACTCCCCCACCGAGACGA 1432 AGTGGGACCTGAGCCAGAATCCCTTCCAAAGCATCGCCTCCTATTCTCCCACCAGCAAGA 1432 ratTas1r2 humanTAS1R2 AATGGGACCGGAGCCAGAATCCCTTCCAGAGCGTCGCCTCCTACTACCCCCTGCAGCGAC 1420 ACTGGAATGGACCTGAATGGACCTTTGAGGTCATTGGTTCTGCCTCACTGTCTCCAGTTC 1426 mouseTas1r1 ratTaslr1 ACTGGAATGGACCTGAATGGACCTTTGAGATCATTGGCTCTGCCTCACTGTCTCCAGTTC 1420 humanTAS1R1 ACTGGAATGGACCCAAGTGGACCTTCACGGTCCTCGGTTCCTCCACATGGTCTCCAGTTC 1423 mouseTas1r3 TGTGGCAGAGCCCTACACCTGTATTACATACTGTGGGCACCT-----TCAACGCA 1450 TGTGGCAGAGCCCTACACCTGTACATACTGTAGGCACCT-----TCAACGGCA 1450 ratTas1r3 catTas1r3 TGTGGCAGGACCCGACCCCGAGCTGCGCACCGTAGGCACCT----TCAAGGGCC 1438 humanTAS1R3 TGTGGCAGGGCTCAGTGCCCAGGCTCCACGACGTGGGCAGGT-----TCAACGGCA 1429 mouseTas1r2 GGCT-GACCTACATTAGCAATGTGTCC--TGGTACACCCCCAACAACACGGTCCCCATAT 1489 GGCT-AACCTACATTAACAATGTGTCC--TGGTACACCCCCAACAACACGGTCCCTGTCT 1489 AGCT-GAAGAACATCCAAGACATCTCC--TGGCACACCGTCAACAACACGGTCCCTATGT 1477 ratTas1r2 humanTAS1R2 mouseTas1r1 ATCTAGACATAAATAAGACAAAAATCCAGTGGCACGGGAAGAACAATCAGGTGCCTGTGT 1486 ratTas1r1 ATCTGGACATAAATAAGACAAAAATCCAGTGGCACGGGAAGAACAATCAGGTGCCTGTGT 1480 humanTAS1R1 AGCTAAACATAAATGAGACCAAAATCCAGTGGCACGGAAAGGACAACCAGGTGCCTAAGT 1483 CCCTTCAGCTGCAGCAGTCTAAAATGTACTGGC-----CAGGCAACCAGGTGCCAGTCT 1504 CCCTTCAGCTGCAGCACTCGAAAATGTATTGGC-----CAGGCAACCAGGTGCCAGTCT 1504 mouseTas1r3 ratTas1r3 catTas1r3 GCCTGGAGCTCTGGCGCTCTCAGATGTGCTGGCACACGCCGGGGAAGCAGCCCGTGT 1498 humanTAS1R3 GCCTCAGGACAGAGCGCCTGAAGATCCGCTGGCACACGTCTGACAACCAGAAGCCCGTGT 1489 mouseTas1r2 CCATGTGTTCTAAGAGTTGCCAGCCTGGGCAAATGAAAAAACCCATAGGCCTCCACCCGT 1549 ratTas1r2 CCATGTGTTCCAAGAGCTGCCAGCCAGGGCAAATGAAAAAGTCTGTGGGCCTCCACCCTT 1549 humanTAS1R2 CCATGTGTTCCAAGAGGTGCCAGTCAGGGCAAAAGAAGAAGCCTGTGGGCATCCACGTCT 1537 mouseTas1r1 CAGTGTGTACCAGGGACTGTCTCGAAGGGCACCACAGGTTGGTCATGGGTTCCCACCACT 1546 ratTas1r1 CAGTGTGTACCACGGACTGTCTGGCAGGGCACCACAGGGTGGTTGTGGGTTCCCACCACT 1540 humanTAS1R1 CTGTGTGTTCCAGCGACTGTCTTGAAGGGCACCAGCGAGTGGTTACGGGTTTCCATCACT 1543 mouseTas1r3 CCCAGTGTTCCCGCCAGTGCAAAGATGGCCAGGTTCGCCGAGTAAAGGGCTTTCATTCCT 1564 ratTaslr3 CCCAGTGCTCCCGGCAGTGCAAAGATGGCCAGGTGCGCAGAGTAAAGGGCTTTCATTCCT 1564 catTas1r3 CCCAGTGCTCCCGGCAGTGCAAGGAAGGCCAGGTGCGCCGCGTGAAGGGCTTCCACTCTT 1558 humanTAS1R3 CCCGGTGCTCGCGGCAGTGCCAGGAGGGCCAGGTGCGCCGGGTCAAGGGGTTCCACTCCT 1549 mouseTas1r2 ratTas1r2 GTTGCTTCGAGTGCTTGGATTGTATGCCAGGCACCTCACCTCAACCGCTCAGCAGATGAGT 1609 humanTAS1R2 GCTGCTTCGAGTGCATCGACTGCCTTCCCGGCACCTTCCTCAACCACACTGAAGATGAAT 1597 mouseTas1r1 GCTGCTTCGAGTGCATGCCCTGTGAAGCTGGGACATTTCTCAAC---ACGAGTGAGCTTC 1603 ratTas1r1 GCTGCTTTGAGTGTGCCCTGCGAAGCTGGGACCTTTCTCAAC---ATGAGTGAGCTTC 1597 GCTGCTTTGAGTGTGCCCTGTGGGGCTGGGACCTTCCTCAAC---AAGAGTGACCTCT 1600 humanTAS1R1 mouseTas1r3 GCTGCTATGACTGCGTGGACTGCAAGGCGGGCAGCTACCGGAAG~--CATCCAGATGACT 1621 ratTas1r3 GCTGCTATGACTGTGTGGACTGCAAGGCAGGGAGCTACCGGAAG---CATCCAGATGACT 1621 catTas1r3 GCTGTTACAACTGCGTGGACTGCAAGGCGGGCAGTTATCAGCGC---AACCCAGATGACC 1615 humanTAS1R3 GCTGCTACGACTGTGGGACTGCGAGGCGGGCAGCTACCGGCAA---AACCCAGACGACA 1606 mouseTas1r2 ratTas1r2 TTAACTGTCTGCCCGGGTTCCATGTGGTCCTACAAGAACGACATCACTTGCTTCC 1669 humanTAS1R2 ATGAATGCCAGGCCTGCCCGAATAACGAGTGGTCCTACCAGAGTGAGACCTCCTGCTTCA 1657 ACACCTGCCAGCCTTGTGGAACAGAAGAATGGGCCCCTGAGGGGAGCTCAGCCTGCTTCT 1663 mouseTas1r1 ACATCTGCCAGCCTTGTGGAACAGAAGAATGGGCACCCAAGGAGGAGCACTACTTGCTTCC 1657 ratTas1r1

ACAGATGCCAGCCTTGTGGGAAAGAAGAGTGGGCACCTGAGGGAAGCCAGACCTGCTTCC 1660

TCACCTGTACTCCATGTAACCAGGACCAGTGGTCCCCAGAGAAAAAGCACAGCCTGCTTAC 1681
TCACCTGTACTCCATGTGGCAAGGATCAGTGGTCCCCAGAAAAAAGCACAACCTGCTTAC 1681

TCCTCTGCACCCAGTGTGACCAGGACCAGTGGTCCCCAGACCGGAGCACACGCTGCTTCG 1675

TCGCCTGCACCTTTTGTGGCCAGGATGAGTGGTCCCCGGAGCGAAGCACACGCTGCTTCC 1666

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Inventors: Xia LI, Weihua LI, Danielle R. REED, Alexander Alexeyevich BACHMANOV,
Joseph G. BRAND
Attomey: Felicity E. Groth
Sheet 6 of 13

Figure 1F

mouseTas1r2	AGCGGCGGCTGGCCTTCCTGGAGTGGCACGAAGTGCCCACTATCGTGGTGACCATCCTGG	1729
ratTas1r2	AGCGGCGGCCTACCTTCCTGGAGTGGCACGAAGTGCCCACCATCGTGGTGGCCATACTGG	1729
humanTAS1R2	AGCGGCAGCTGGTCTTCCTGGAATGGCATGAGGCACCCACC	1717
mouseTas1r1	CACGCACCGTGGAGTTCTTGGGGTGGCATGAACCCATCTCTTTGGTGCTATTAGCAGCTA	
ratTas1r1	CACGCACGGTGGAGTTCTTGGCTTGGCATGAACCCATCTCTTTGGTGCTAATAGCAGCTA	
humanTAS1R1	CGCGCACTGTGGTGTTTTTGGCTTTGCGTGAGCACACCTCTTGGGTGCTGCTGGCAGCTA	
mouseTas1r3	CTCGCAGGCCCAAGTTTCTGGCTTGGGGGGGGCCAGTTGTGCTGTCACTCCTCCTGCTGC	1741
ratTas1r3	CTCGCAGGCCCAAGTTTCTGGCTTGGGGGGGGCCAGCTGTGCTGTCACTTCTCCTGCTGC	1741
catTas1r3	CCCGCAAGCCCATGTTCCTGGCATGGGGGGAGCCAGCTGTGCTGCTACTGCTCGCGCTGC	1735
humanTAS1R3	GCCGCAGGTCTCGGTTCCTGGCATGGGGCGAGCCGGCTGTGCTGCTGCTGCTGCTGC	1726
	**	
mouseTas1r2	CCGCCCTGG-GCTTCATCAGTACGCTGGCCATTCTGCTCATCTTCTGGAGACATTTCCAG	1700
ratTas1r2	CTGCCCTGG-GCTTCTTCAGTACACTGGCCATTCTTTTCATCTTCTGGAGACATTTCCAG	
humanTAS1R2	CCGCCCTGG-GCTTCCTCAGCACCCTGGCCATCCTGGTGATATTCTGGAGGCACTTCCAG	
mouseTaslrl	ACACGCTATTGCTGCTGCTGATTGGGACTGCTGGCC-TGTTTGCCTGGCGTCTTCAC	1782
ratTaslr1	ACACGCTATTGCTGCTGCTGGTTGGGACTGCTGGCC-TGTTTGCCTGGCATTTTCAC	1776
humanTAS1R1	ACACGCTGCTGCTGCTGCTTGGGACTGCTGGCC-TGTTTGCCTGGCACCTAGAC	1779
mouseTas1r3	TTTGCCTGGTGCTGGGTCTA-GCACTGGCTGCTCTGGGGCTCTCTGTCCACCACTGGGAC	1800
ratTaslr3	TTTGCCTGGTGCTGGGCCTG-ACACTGGCTGCCCTGGGGCTCTTTGTCCACTACTGGGAC	1800
catTaslr3	TGGCTCTGGCGCTGGCCTG-GCGCTGGCAGCCCTGGGGCTCTTCCTCTGGCACTCGGAC	
humanTAS1R3	TGAGCCTGGCGCTGGCCTT-GTGCTGGCTGCTTTGGGGCTGTTCGTTC	
Hamaning	** *** * *** * * * * * * * * * * * * *	1,00
mouseTas1r2	ACGCCCATGGTGCGCTCGGCGGCCGCCCCATGTGCTTCCTGATGCTGGTGCCCCTGCTG	
ratTas1r2	ACACCCATGGTGCGCTCGGCCGGTGGCCCCATGTGCTTCCTGATGCTCGTGCCCCTGCTG	1848
humanTAS1R2	ACACCCATAGTTCGCTCGGCTGGGGGCCCCATGTGCTTCCTGATGCTGACACTGCTGCTG	1836
mouseTas1r1	ACGCCTGTTGTGAGGTCAGCTGGGGTAGGCTGTGCTTCCTCATGCTGGGTTCCTTGGTA	1842
ratTaslrl	ACACCTGTAGTGAGGTCAGCTGGGGGTAGGCTGTGCTTCCTCATGCTGGGTTCCCTGGTG	1836
humanTAS1R1	ACCCCTGTGGTGAGGTCAGCAGGGGGCCGCCTGTGCTTTCTTATGCTGGGCTCCCTGGCA	
mouseTas1r3	AGCCCTCTTGTCCAGGCCTCAGGTGGCTCACAGTTCTGCTTTGGCCTGATCTGCCTAGGC	
ratTaslr3	AGCCCTCTTGTTCAGGCCTCAGGTGGGTCACTGTTCTGCTTTGGCCTGATCTGCCTAGGC	
catTas1r3	AGCCCGCTGGTTCAGGCCTCAGGTGGGCCACGGGCCTGCTTTGGCCTGGCTTGCCTGGGC	
humanTAS1R3	AGCCCACTGGTTCAGGCCTCGGGGGGGCCCCTGGCCTGCTTTGGCCTGGTGTGCCTGGGC	1845
	* ** * * * * * * * * * * * * *	
mouseTas1r2	CTGGCGTTCGGGATGGTCCCCGTGTATGTGGGCCCCCCCACGGTCTTCTCCTGTTTCTGC	1908
ratTas1r2	CTGGCGTTTGGGATGGTGCCCGTGTATGTGGGGCCCCCCACGGTCTTCTCATGCTTCTGC	
humanTAS1R2	GTGGCATACATGGTGGTCCCGGTGTACGTGGGGCCCCCAAGGTCTCCACCTGCCTCTGC	
mouseTas1r1	GCTGGGAGTTGCAGCCTCTACAGCTTCTTCGGGAAGCCCACGGTGCCCGCGTGCTTGCT	
ratTas1r1	GCCGGAAGTTGCAGCTTCTATAGCTTCTTCGGGGAGCCCACGGTGCCCGCGTGCTTGCT	
humanTAS1R1	GCAGGTAGTGGCAGCCTCTATGGCTTCTTTGGGGAACCCACAAGGCCTGCGTGCTTGCT	1899
mouseTaslr3	CTCTTCTGCCTCAGTGTCCTTCTGTTCCCAGGGCGGCCAAGCTCTGCCAGCTGCCTTGCA	1920
ratTaslr3	CTCTTCTGCCTCAGTGTCCTTCTGTTCCCAGGACGACCACGCTCTGCCAGCTGCCTTGCC	1920
catTaslr3	CTGGTCTGCCTCAGTGTCCTGTTCCCTGGCCAGCCAGGCCCTGCCAGCTGCCTGGCC	1914
humanTAS1R3	CTGGTCTGCCTCAGCGTCCTCCTGTTCCCTGGCCAGCCCAGCCCTGCCCGATGCCTGGCC	
	* * ** ** **	
	CGCCAGGCTTTCTTCACCGTTTGCTTCTCCGTCTGCCTCTCCTGCATCACGGTGCGCTCC	1000
mouseTas1r2		
ratTaslr2	CGACAGGCTTTCTTCACCGTCTGCTTCTCCATCTGCCTATCCTGCATCACCGTGCGCTCC	
humanTAS1R2	CGCCAGGCCCTCTTTCCCCTCTGCTTCACAATTTGCATCTCCTGTATCGCCGTGCGTTCT	
mouseTas1r1	CGTCAGCCCCTCTTTTCTCTCGGGTTTGCCATTTTCCTCTCTGTCTG	1962
ratTas1r1	CGTCAGCCCCTCTTTCCTCGGGTTTGCCATCTTCCTCTCCTGCCTG	1956
humanTAS1R1	CGCCAGGCCCTCTTTGCCCTTGGTTTCACCATCTTCCTGTCCTGCCTG	1959
mouseTas1r3	CAACAACCAATGGCTCACCTCCCTCTCACAGGCTGCCTGAGCACCACTCTTCCTGCAAGCA	1980
ratTas1r3	CAACAACCAATGGCTCACCTCCCTCTCACAGGCTGCCTGAGCACACTCTTCCTGCAAGCA	
catTaslr3	CAGCAGCCACTGTTCCACCTCCCACTCACTGGCTGCCTGAGCACGTTTTTCCTGCAAGCG	
humanTAS1R3	CAGCAGCCCTTGTCCCACCTCCCGCTCACGGGCTGCCTGAGCACACTCTTCCTGCAGGCG  * ** * * * * * * * * * * * * * * * *	1202
mouseTas1r2	TTCCAGATTGTGTGCGTCTTCAAGATGGCCAGACGCCTGCCAAGCGCCTACGGTTTCTGG	
ratTas1r2	TTCCAGATCGTGTGTCTTCAAGATGGCCAGACGCCTGCCAAGTGCCTACAGTTTTTGG	2028
humanTAS1R2	TTCCAGATCGTCTGCGCCTTCAAGATGGCCAGCCGCTTCCCACGCGCCTACAGCTACTGG	2016
mouseTas1r1	TTCCAACTGGTCATCATCTTCAAGTTTTCTACCAAGGTACCCACATTCTACCACACTTGG	
ratTas1r1	TTCCAACTGGTCATCATCTTCAAGTTTTCTACCAAGGTGCCCACATTCTACCGTACCTGG	
humanTAS1R1	TTCCAACTAATCATCTTCAAGTTTTCCACCAAGGTACCTACATTCTACCACGCCTGG	
mouseTaslr3		
	GCTGAGACCTTTGTGGAGTCTGAGCTGCCACTGAGCTGGGCAAACTGGCTATGCAGCTAC	
ratTas1r3	GCCGAGATCTTTGTGGAGTCTGAGCTGCCACTGAGTTGGGCAAACTGGCTCTGCAGCTAC	
catTas1r3	GCCGAGATATTTGTGGGGTCGGAGCTGCCACCAAGCTGGGCTGAGAAGATGCGTGGCCGC	
humanTAS1R3	GCCGAGATCTTCGTGGAGTCAGAACTGCCTCTGAGCTGGGCAGACCGGCTGAGTGGCTGC	2025
	* * * * * *	

Docket No.: MON-3042

App No.: Not Yet Assigned

Title: TASTE RECEPTOR OF THE TIR FAMILY FROM DOMESTIC CAT Inventors: Xia LI, Weihua LI, Danielle R. REED, Alexander Alexeyevich BACHMANOV, Joseph G. BRAND

Attorney: Felicity E. Groth

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Figure 1G

rigule 10		
mouseTas1r2	ATGCGTTACCACGGGCCCTACGTCTTTGTGGCCTTCATCACGGCCGTCAAGGTGGCCCT-	2087
ratTas1r2	ATGCGTTACCACGGGCCCTATGTCTTCGTGGCCTTCATCACGGCCATCAAGGTGGCCCT-	2087
humanTAS1R2	GTCCGCTACCAGGGGCCCTACGTCTCTATGGCATTTATCACGGTACTCAAAATGGTCAT-	2075
mouseTaslr1	GCCCAAAACCATGGTGCCGGAATATTCGTCATTGTCAGCTCCACGGTCCATTTGTTCCTC	
ratTaslrl	GCCCAAAACCATGGTGCAGGTCTATTCGTCATTGTCAGCTCCACGGTCCATTTGCTCATC	
humanTAS1R1	GTCCAAAACCACGGTGCTGGCCTGTTTGTGATGATCAGCTCAGCGGCCCAGCTGCTTATC	
mouseTas1r3	CTTCGGGGACTCTGGCCTAGTGGTACTGTTGGCCACTTTTGTGGAGGCAGCACTA	
ratTaslr3	CTTCGGGGCCCCTGGGCTTGGCTGGTACTGCTGGCCACTCTTGTGGAGGCTGCACTA	
catTas1r3	CTGCGGGGGCCCTGGCCTGGCTGGTGCTGCTTGCTATGCTGGCAGAAGCCGCATTG	
humanTAS1R3	CTGCGGGGGCCCTGGCTGGTGGTGCTGCTGGCCATGCTGGTGGAGGTCGCACTG	2085
	* * * * * * * * *	
mouseTas1r2	GGTGGCAGGCAACATGCTGGCCACCACCATCAACCCCATTGGCCGGACCGACC	
ratTas1r2	GGTGGTGGCCAACATGCTGGCCACCACCATCAACCCCATTGGCCGGACCGGACCCGGATGA	
humanTAS1R2	TGTGGTAATTGGCATGCCACGGGCCTCAGTCCCACCACCGTACTGACCCCGATGA	
mouseTaslrl ratTaslrl	TGTCTCACGTGGCTTGCAATGTGGACCCCACGGCCCACGGGAGTACCAGCGCTT	
	TGTCTCACATGGCTTGTAATGTGGACCCCACGACCCACCAGGGAATACCAGCGCTT	
humanTAS1R1	TGTCTAACTTGGCTGGTGGTGTGGACCCCACTGCCTGCTAGGGAATACCAGCGCTT TGTGCCTGGTATTTGATCGCTTTCCCACCAGAGGTGGTGACAGACTGGTCAGTGCTG	
mouseTas1r3 ratTas1r3		
catTas1r3	TGTGCCTGGTACTTGATGGCTTTCCCTCCAGAGGTGGTGACAGATTGGCAGGTGCTG TGTGCCTGGTACCTGGTAGCCTTCCCGCCAGAGGTGGTGACGGACTGGCGGGTACTG	
humanTAS1R3	TGCACCTGGTACCTGGTGGCCTTCCCGCCGGAGGTGGTGACGGACTGGCACATGCTG	
namaninoino	* * *	2142
mouseTas1r2	CCCCAATATCATAATCCTCTCCTGCCACCCTAACTACCGCAACGGGCTACTCTTCAACAC	2207
ratTas1r2	CCCCAACATCATGATCCTCTCGTGCCACCCTAACTACCGCAACGGGCTACTGTTCAACAC	2207
humanTAS1R2	CCCCAAGATCACAATTGTCTCCTGTAACCCCAACTACCGCAACAGCCTGCTGTTCAACAC	
mouseTas1r1	CCCCCATCTGGTGATTCTTGAGTGC-ACAGAGGTCAACTCTGTGGGCTTCCTGGTGGCTT	
ratTas1r1	CCCCCATCTGGTGATTCTCGAGTGC-ACAGAGGTCAACTCTGTAGGCTTCCTGTTGGCTT	
humanTAS1R1	CCCCCATCTGGTGATGCTTGAGTGC-ACAGAGACCAACTCCCTGGGCTTCATACTGGCCT	2194
mouseTas1r3	CCCACAGA-GGTACTGGAGCACTGCCACGTGCGTT-CCTGGGTCAGCCTGGGCTTGGTGC	
ratTas1r3	CCCACGGA-GGTACTGGAACACTGCCGCATGCGTT-CCTGGGTCAGCCTGGGCTTGGTGC	
catTaslr3	CCCACAGA-GGCGCTGGTGCACTGCCACGTGCACT-CCTGGATCAGCTTCGGCCTGGTGC	
humanTAS1R3	CCCACGGA-GGCGCTGGTGCACTGCCGCACACGCT-CCTGGGTCAGCTTCGGCCTAGCGC	2200
	***	
mouseTas1r2	CAGCATG-GACTTGCTGCTGTCCGTGCTGGGTTTCAGCTTCGCGTACGTGGGCAAGGAAC	2266
ratTas1r2		2266
humanTAS1R2	CAGCCTG-GACCTGCTCTCAGTGGTGGGTTTCAGCTTCGCCTACATGGGCAAAGAGC	
mouseTas1r1	TCGCACACAACATCCTCCTCCCATCAGCACCTTTGTCTGCAGCTACCTGGGTAAGGAAC	
ratTas1r1	TCACCCACAACATTCTCCTCTCCATCAGTACCTTCGTCTGCAGCTACCTGGGTAAGGAAC	
humanTAS1R1	TCCTCTACAATGGCCTCCTCTCCATCAGTGCCTTTGCCTGCAGCTACCTGGGTAAGGACT	2254
mouseTas1r3	ACATCACCAATGCAATGTTAGCTTTCCTCTGCTTTCTGGGCACTTTCCTGGTACAGAGCC	
ratTas1r3	ACATCACCAATGCAGTGTTAGCTTTCCTCTGCTTTCTGGGCACTTTCCTGGTACAGAGCC	2275
catTas1r3	ATGCCACTAACGCCATGCTGGCCTTCCTCTGCTTCCTGGGCACTTTCCTGGTGCAGAGCC	2269
humanTAS1R3	ACGCCACCAATGCCACGCTGGCCTTTCTCTGCTTCCTGGGCACTTTCCTGGTGCGGAGCC	2260
	* * * * * * * * * * * * * * * * * * * *	
	######################################	2226
mouseTas1r2	TGCCCACCAACTACAACGAAGCCAAGTTCATCACCCTCAGCATGACCTTCTCCTTCACCT	2326
ratTas1r2 humanTAS1R2	TGCCCACCAACTACAACGAAGCCAAGTTCATCACTCTCAGCATGACCTTCTCCTTCACCT TGCCCACCAACTACAACGAGGCCAAGTTCATCACCCTCAGCATGACCTTCTATTTCACCT	2326
mouseTas1r1	TGCCGGAGAACTATAACGAAGCCAAATGTGTCACCTTCAGCCTGCTCCTCCACTTCGTAT	
ratTaslrl	TGCCAGAGAACTATAATGAAGCCAAATGTGTCACCTTCAGCCTGCTCCTCCACTTCGTAT	
humanTAS1R1	TGCCAGAGAACTACAACGAGGCCAAATGTGTCACCTTCAGCCTGCTCTTCAACTTCGTGT	
mouseTas1r3	AGCCTGGCCGCTACAACCGTGCCCGTGGTCTCACCTTCGCCATGCTAGCTTATTTCATCA	
ratTas1r3	AGCCTGGTCGCTATAACCGTGCCCGTGGCCTCACCTTCGCCATGCTAGCTTATTTCATCA	
catTas1r3	GGCCAGGCCGCTACAATGGTGCCCGCGGCCTCACCTTTGCCATGCTGGCCTACTTCATCA	
humanTAS1R3	AGCCGGGCCGCTACAACCGTGCCCTGGCCTCACCTTTGCCATGCTGGCCTACTTCATCA	
	*** *** *** *** * * * * * * *	
mouseTas1r2	CCTCCATCTCCCTCTGCACGTTCATGTCTGTCCACGATGGCGTGCTGGTCACCATCATGG	
ratTas1r2	CCTCCATCTCCCTCTGCACCTTCATGTCTGTGCACGACGGCGTGCTGGTCACCATCATGG	
humanTAS1R2	CATCCGTCTCCCTCTGCACCTTCATGTCTGCCTACAGCGGGGTGCTGGTCACCATCGTGG	
mouseTaslrl	CCTGGATCGCTTTCTTCACCATGTCCAGCATTTACCAGGGCA-GCTACCTACCCGCGG	
ratTaslrl	CCTGGATCGCCTTCTTCACCATGGCCAGCATTTACCAGGGCA-GCTACCTGCCTGCGG	
humanTAS1R1	CCTGGATCGCCTTCTTCACCACGGCCAGCGTCTACGACGGCA-AGTACCTGCCTGCGG	
mouseTaslr3	CCTGGGTCTCTTTTGTGCCCCTCCTGGCCA-ATGTGCAGGTGGCCTACCAGCCAGCTG	
ratTas1r3 catTas1r3	TCTGGGTCTCTTTTGTGCCCCTCCTGGCTA-ATGTGCAGGTGGCCTACCAGCCAGCTG CCTGGATCTCCTTTGTGCCCCTCTTTGCCA-ATGTGCACGTGGCCTACCAGCCTGCCG	
humanTAS1R3	CCTGGGTCTCCTTTGTGCCCCTCTTTGCCA-ATGTGCACGTGGCCTACCAGCCTGCCG CCTGGGTCTCCTTTGTGCCCCTCCTGGCCA-ATGTGCAGGTGGTCCTCAGGCCCGCCG	
	* ** * * * * * * * * * * * * * * * * *	2311

Docket No.: MON-3042

App No.: Not Yet Assigned Filed: Herewith

Title: TASTE RECEPTOR OF THE TIR FAMILY FROM DOMESTIC CAT
Inventors: Xia LI, Weihua LI, Danielle R. REED, Alexander Alexeyevich BACHMANOV,
Joseph G. BRAND

Attorney: Felicity E. Groth
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## Figure 1H

mouseTas1r2	ATCTCCTGGTCACTGTGCTCAACTTTCTGGCCATCGGCTTGGGGTACTTTGGCCCCA	2443
ratTaslr2	ACCTCCTGGTCACTGTGCTCAACTTCCTGGCCATCGGCTTGGGATACTTTGGCCCCA	2443
humanTAS1R2	ACCTCTTGGTCACTGTGCTCAACCTCCTGGCCATCAGCCTGGGCTACTTCGGCCCCA	2431
mouseTas1r1	TCAATGTGCTGGCAGGGCTGGCCACTCTGAGTGGCGGCTTCAGCGGCTATTTCCTCCCTA	2434
ratTaslrl	TCAATGTGCTGGCAGGGCTGACCACACTGAGCGGCGGCTTCAGCGGTTACTTCCTCCCCA	2428
humanTAS1R1	CCAACATGATGGCTGGGCTGAGCAGCCTGAGCAGCGGCTTCGGTGGGTATTTTCTGCCTA	2431
mouseTas1r3	TGCAGATGGGTGCTATCCTAGTCTGTGCCCTGGGCATCCTGGTCACCTTCCACCTGCCCA	2452
ratTas1r3	TGCAGATGGGTGCTATCTTATTCTGTGCCCTGGGCATCCTGGCCACCTTCCACCTGCCCA	2452
catTaslr3	TGCAGATGGGCACCATCCTCTGTGCCCTGGGTATCCTAGCCACCTTCCACCTGCCCA	2446
humanTAS1R3	TGCAGATGGGCGCCCTCCTGCTCTGTGTCCTGGGCATCCTGGCTGCCTTCCACCTGCCCA	2437
	** * * * * * * * *	
mouseTas1r2	AGTGTTACATGATCCTTTTCTACCCGGAGCGCAACACTTCAGCTTATTTCAATAGCATGA	2503
ratTas1r2	AGTGTTACATGATCCTTTTCTACCCGGAGCGCAACACCTCAGCCTATTTCAATAGCATGA	2503
humanTAS1R2	AGTGCTACATGATCCTCTTCTACCCGGAGCGCAACACGCCCGCC	2491
mouseTas1r1	AATGCTACGTGATTCTCTGCCGTCCAGAACTCAACAACACAGAACACTTTCAGGCCTCCA	2494
ratTaslrl	AGTGCTATGTGATTCTCTGCCGTCCAGAACTCAACAATACAGAACACTTTCAGGCCTCCA	2488
humanTAS1R1	AGTGCTACGTGATCCTCTGCCGCCCAGACCTCAACAGCACAGAGCACTTCCAGGCCTCCA	2491
mouseTas1r3	AGTGCTATGTGCTTCTTTGGCTGCCAAAGCTCAACACCCCAGGAGTTCTTCCTGGGAAGGA	2512
ratTas1r3	AATGCTATGTACTTCTGTGGCTGCCAGAGCTCAACACCCCAGGAGTTCTTCCTGGGAAGGA	2512
catTas1r3	AGTGCTACCTGCTGCAGCGGCCGGAGCTCAACACCCCCTGAGTTCTTCCTGGAAGACA	2506
humanTAS1R3	GGTGTTACCTGCTCATGCGGCAGCCAGGGCTCAACACCCCCGAGTTCTTCCTGGGAGGGG	2497
	** ** * * * * * ***** * **	
mouseTaslr2	TTCAGGGCTACACGATGAGGAAGAGCTAG	2532
ratTaslr2	TCCAGGGCTACACCATGAGGAAGAGC	2529
humanTAS1R2	TCCAGGGCTACACCATGAGGAGGGACTAG	2520
mouseTaslrl	TCCAGGACTACACGAGGCGCTGCGGCACTACCTGA	
ratTas1r1	TCCAGGACTACACGAGGCGCTGCGGCACTACC	2520
humanTAS1R1	TTCAGGACTACACGAGGCGCTGCGGCTCCACCTGA	2526
mouseTas1r3	ATGCCAAGAAAGCAGCAGATGAGAAC-AGTGGCGGTGGTGAGGCAGCTCAGGGACACAAT	2571
ratTaslr3	GCCCCAAGGAAGCATCAGATGGGAAT-AGTGGTAGTAGTGAGGCAACTCGGGGACACAGT	2571
catTas1r3	ATGCCAGAGCACAGGGCAGCAGTTGGGGGCAGGGGGGGGGAGAATCGGGGCAAAAAC	
humanTAS1R3	GCCCTGGGGATGCCCAAGGCCAGAATGACGGGAACACAGGAAATCAGGGGAAACAT	2553
	*	
mouseTas1r2		
ratTas1r2		
humanTAS1R2		
mouseTas1r1		
ratTas1r1		
humanTAS1R1		
mouseTaslr3		
	GAATGA 2577	
ratTas1r3	GAATGA 2577	
ratTas1r3 catTas1r3	GAATGA 2577 AAGTGA 2569	
ratTas1r3	GAATGA 2577	

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Inventors: Xia LI, Weihua LI, Danielle R. REED, Alexander Alexeyevich BACHMANOV,

Joseph G. BRAND

Attorney: Felicity E. Groth Phone: (215) 568-3100
Sheet 9 of 13

Figure 2A CLUSTAL W (1.82) multiple amin acid sequence alignment f T1Rs:

```
MLGPAVLGLS----LWALLHPGTGAPLCLSQQLRMKGDYVLGGLFPLGEAEEAGLRSR-- 54
humanT1R3
catT1R3
                MPGLALLGLTALLDHGEGATSCLSQQLRMQGDYVLGGLFPLGSAEGTGLGDG-- 58
mouseT1R3
                MPALAIMGLS----LAAFLELGMGASLCLSQQFKAQGDYILGGLFPLGSTEEATLNQR-- 54
                MPGLAILGLS----LAAFLELGMGSSLCLSQQFKAQGDYILGGLFPLGTTEEATLNQR-- 54
ratT1R3
mouseT1R1
                MLFWAAHLLLSLQLAVAYCWAFSCQRTESSPGFSLPGDFLLAGLFSLHADCLQVRHR--P 58
ratT1R1
                MLFWAAHLLLSLQL--VYCWAFSCQRTESSPGFSLPGDFLLAGLFSLHGDCLQVRHR--P 56
                MLLCTARLVG-LQLLISCCWAFACHSTESSPDFTLPGDYLLAGLFPLHSGCLQVRHR--P 57
humanT1R1
mouseT1R2
                --MGPQARTLHLLFLLLHALPKPVMLVGNSD-FHLAGDYLLGGLFTLHANVKSVSHLSYL 57
ratT1R2
                 --MGPQARTLCLLSLLLHVLPKPGKLVENSD-FHLAGDYLLGGLFTLHANVKSISHLSYL 57
humanT1R2
                 --MGPRAKTICSLFFLLWVLAEP---AENSD-FYLPGDYLLGGLFSLHANMKGIVHLNFL 54
                                                     **::*.***.*
                        (possible functional amino acid substitution)
                TRPSSPVCTRFSSNGLLWALAMKMAVEEINNKSDLLPGLRLGYDLFDTCSEPVVAMKPSL 114
humanT1R3
catT1R3
                LOPNATVCTRFSSLGLLWALAVKMAVEEINNGSALLPGLHLGYDLFDTCSEPMVAMKPSL 118
mouseT1R3
                TOPNS PCNRFSPLGLFLAMAMKMAVEEINNGSALLPGLRLGYDLFDTCSEPVVTMKSSL 114
                TOPNGILCTRFSPLGLFLAMAMKMAVEEINNGSALLPGLRLGYDLFDTCSEPVVTMKPSL 114
ratT1R3
                LVTSCDRSDSFNGHGYHLFQAMRFTVEEINNSTALLPNITLGYELYDVCSE-SSNVYATL 117
mouseT1R1
                LVTSCDRPDSFNGHGYHLFQAMRFTVEEINNSSALLPNITLGYELYDVCSE-SANVYATL 115
ratT1R1
                EVTLCDRSCSFNEHGYHLFQAMRLGVEEINNSTALLPNITLGYQLYDVCSD-SANVYATL 116
humanT1R1
mouseT1R2
                QVPKCN-EYNMKVLGYNLMQAMRFAVEEINNCSSLLPGVLLGYEMVDVCYL-SNNIQPGL 115
ratT1R2
                QVPKCN-EFTMKVLGYNLMQAMRFAVEEINNCSSLLPGVLLGYEMVDVCYL-SNNIHPGL 115
humanT1R2
                QVPMCK-EYEVKVIGYNLMQAMRFAVEEINNDSSLLPGVLLGYEIVDVCYI-SNNVQPVL 112
                                    *::: ***** : ***.: ***:: *.*
humanT1R3
                MFLAKAGSRDIAAYCNYTQYQPRVLAVIGPHSSELAMVTGKFFSFFLMPQVSYGASMELL 174
                VFMAKAGSCSIAAYCNYTQYQPRVLAVIGPHSSELALVTGKFFSFFLVPQVSYGASTDRL 178
catT1R3
mouseT1R3
                MFLAKVGSQSIAAYCNYTQYQPRVLAVIGPHSSELALITGKFFSFFLMPQVSYSASMDRL 174
ratT1R3
                MFMAKVGSQSIAAYCNYTQYQPRVLAVIGPHSSELALITGKFFSFFLMPQVSYSASMDRL 174
mouseT1R1
                RVLAQQGTGHLEMQRDLRNHSSKVVALIGPDNTDHAVTTAALLSPFLMPLVSYEASSVIL 177
ratT1R1
                RVLALQGPRHIEIQKDLRNHSSKVVAFIGPDNTDHAVTTAALLGPFLMPLVSYEASSVVL 175
humanT1R1
                RVLSLPGQHHIELQGDLLHYSPTVLAVIGPDSTNRAATTAALLSPFLVPMISYAASSETL 176
                YFLSQI-DDFLPILKDYSQYRPQVVAVIGPDNSESAITVSNILSYFLVPQVTYSAITDKL 174
mouseT1R2
ratT1R2
                YFLAQD-DDLLPILKDYSQYMPHVVAVIGPDNSESAITVSNILSHFLIPQITYSAISDKL 174
                YFLAHE-DNLLPIQEDYSNYISRVVAVIGPDNSESVMTVANFLSLFLLPQITYSAISDEL 171
humanT1R2
                               : :: . *:*.***..:: .
humanT1R3
                SARETFPSFFRTVPSDRVQLTAAAELLQEFGWNWVAALGSDDEYGRQGLSIFSALAAAR- 233
catT1R3
                SNREIFPSFFRTVPSDQVQVAAMVELLEELGWNWVAAVGSDDEYGRQGLSLFSGLASAR- 237
mouseT1R3
                SDRETFPSFFRTVPSDRVQLQAVVTLLQNFSWNWVAALGSDDDYGREGLSIFSSLANAR- 233
ratT1R3
                SDRETFPSFFRTVPSDRVQLQAVVTLLQNFSWNWVAALGSDDDYGREGLSIFSGLANSR- 233
                SGKRKFPSFLRTIPSDKYQVEVIVRLLQSFGWVWISLVGSYGDYGQLGVQALEELATPR- 236
mouseT1R1
                SAKRKFPSFLRTVPSDRHQVEVMVQLLQSFGWVWISLIGSYGDYGQLGVQALEELAVPR- 234
ratT1R1
                SVKRQYPSFLRTIPNDKYQVETMVLLLQKFGWTWISLVGSSDDYGQLGVQALENQATGQ- 235
humanT1R1
mouseT1R2
                RDKRRFPAMLRTVPSATHHIEAMVQLMVHFQWNWIVVLVSDDDYGRENSHLLSQRLTNTG 234
ratT1R2
                RDKRHFPSMLRTVPSATHHIEAMVQLMVHFQWNWIVVLVSDDDYGRENSHLLSQRLTKTS 234
humanT1R2
                RDKVRFPALLRTTPSADHHVEAMVQLMLHFRWNWIIVLVSSDTYGRDNGQLLGERVARR- 230
                                 :: . . *: : * *: : * . **: .
                GICIAHEGLVPLPRADDSR----LGKVQDVLHQVNQSSVQVVLLFASVHAAHALFNYSIS 289
humanT1R3
                GICIAHEGLVPLP-PGSLR----LGALQGLLRQVNQSSVQVVVLFSSAHAARTLFSYSIR 292
catT1R3
                GICIAHEGLVPQHDTSGQQ----LGKVLDVLRQVNQSKVQVVVLFASARAVYSLFSYSIH 289
mouseT1R3
                GICIAHEGLVPQHDTSGQQ----LGKVVDVLRQVNQSKVQVVVLFASARAVYSLFSYSIL 289
ratT1R3
mouseT1R1
                GICVAFKDVVPLSAQAG-----DPRMQRMMLRLARARTTVVVVFSNRHLAGVFFRSVVL 290
                GICVAFKDIVPFSARVG-----DPRMQSMMQHLAQARTTVVVVFSNRHLARVFFRSVVL 288
ratT1R1
                GICIAFKDIMPFSAQVG-----DERMQCLMRHLAQAGATVVVVFSSRQLARVFFESVVL 289
humanT1R1
mouseT1R2
                DICIAFQEVLPVPEPNQAVRPEEQDQLDNILDKLRRTSARVVVIFSPELSLHNFFREVLR 294
                DICIAFQEVLPIPESSQVMRSEEQRQLDNILDKLRRTSARVVVVFSPELSLYSFFHEVLR 294
ratT1R2
humanT1R2
                DICIAFQETLPTLQPNQNMTSEERQRLVTIVDKLQQSTARVVVVFSPDLTLYHFFNEVLR 290
                .**:*.: :*
                                          : :: :: :: . **::*:
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App No.: Not Yet Assigned Filed: Herewith Title: TASTE RECEPTOR OF THE TIR FAMILY FROM DOMESTIC CAT Inventors: Xia LI, Weihua LI, Danielle R. REED, Alexander Alexeyevich BACHMANOV, Joseph G. BRAND Attorney: Felicity E. Groth Phone: (215) 568-3100

Attorney: Felicity E. Groth Ph Sheet 10 of 13

Figure 2B

```
SRLSPKVWVASEAWLTSDLVMGLPGMAQMGTVLGFLQRGAQLHEFPQYVKTHLALATDPA 349
humanT1R3
                CKLSPKVWVASEAWLTSDLVMTLPGMPGVGTVLGFLQQGAPMPEFPSYVRTRLALAADPA 352
catT1R3
                HGLSPKVWVASESWLTSDLVMTLPNIARVGTVLGFLQRGALLPEFSHYVETHLALAADPA 349
mouseT1R3
                HDLSPKVWVASESWLTSDLVMTLPNIARVGTVLGFLQRGALLPEFSHYVETRLALAADPT 349
ratT1R3
mouseT1R1
                ANLTGKVWIASEDWAISTYITNVPGIQGIGTVLGVAIQQRQVPGLKEFEESYVQAVMGAP 350
ratT1R1
                ANLTGKVWVASEDWAISTYITSVTGIQGIGTVLGVAVQQRQVPGLKEFEESYVRAVTAAP 348
humanT1R1
                TNLTGKVWVASEAWALSRHITGVPGIQRIGMVLGVAIQKRAVPGLKAFEEAYARADKKAP 349
mouseT1R2
                WNFTGFVWIASESWAIDPVLHNLTELRHTGTFLGVTIQRVSIPGFSQFRVRHDKPEYPMP 354
ratT1R2
                WNFTGFVWIASESWAIDPVLHNLTELRHTGTFLGVTIQRVSIPGFSQFRVRRDKPGYPVP 354
                QNFTGAVWIASESWAIDPVLHNLTELGHLGTFLGITIQSVPIPGFSEFREWGPQAGPPPL 350
humanT1R2
                                . : :. :
                                              * .**.
humanT1R3
                FCSALGEREQGLEEDVVGQRCPQCDCITLQNVS-----AGLNHHQTFSVYAAVYSVA 401
                FCASLDAEQPGLEEHVVGPRCPQCDHVTLENLS-----AGLLHHQTFAAYAAVYGVA 404
catT1R3
mouseT1R3
                FCASLN-AELDLEEHVMGQRCPRCDDIMLQNLSSGLLQNLSAGQLHHQIFATYAAVYSVA 408
                FCASLK-AELDLEERVMGPRCSQCDYIMLQNLSSGLMQNLSAGQLHHQIFATYAAVYSVA 408
ratT1R3
                RTCPEG-----SWCGTNQLCRECHAFTTWNMP-----ELGAFSMSAAYNVYEAVYAVA 398
mouseT1R1
                SACPEG-----SWCSTNQLCRECHTFTTRNMP-----TLGAFSMSAAYRVYEAVYAVA 396
ratT1R1
                RPCHKG----SWCSSNQLCRECQAFMAHTMP-----KLKAFSMSSAYNAYRAVYAVA 397
humanT1R1
                NETSLR----TTC-NQDCDACMNITESFNN-----VLMLSGERVVYSVYSAVYAVA 400
NTTNLR-----TTC-NQDCDACLNTTKSFNN-----ILILSGERVVYSVYSAVYAVA 400
mouseT1R2
ratT1R2
humanT1R2
                SRTSQS-----YTC--NQECDNCLNATLSFNT-----ILRLSGERVVYSVYSAVYAVA 396
                QALHNTLQCNASGCPAQDPVKPWQLLENMYNLTFHVGGLPLRFDSSGNVDMEYDLKLWVW 461
humanT1R3
                QALHNTLRCNASGCPRREPVRPWQLLENMYNVSFRARGLALQFDASGNVNVDYDLKLWVW 464
catT1R3
mouseT1R3
                QALHNTLQCNVSHCHVSEHVLPWQLLENMYNMSFHARDLTLQFDAEGNVDMEYDLKMWVW 468
ratT1R3
                QALHNTLQCNVSHCHTSEPVQPWQLLENMYNMSFRARDLTLQFDAKGSVDMEYDLKMWVW 468
mouseT1R1
                HGLHQLLGCTSGTCA-RGPVYPWQLLQQIYKVNFLLHKKTVAFDDKGDPLGYYDIIAWDW 457
                HGLHQLLGCTSEICS-RGPVYPWQLLQQIYKVNFLLHENTVAFDDNGDTLGYYDIIAWDW 455
ratT1R1
humanT1R1
                HGLHQLLGCASGACS-RGRVYPWQLLEQIHKVHFLLHKDTVAFNDNRDPLSSYNIIAWDW 456
                HTLHRLLHCNQVRCT-KQIVYPWQLLREIWHVNFTLLGNQLFFDEQGDMPMLLDIIQWQW 459
mouseT1R2
                HALHRLLGCNRVRCT-KQKVYPWQLLREIWHVNFTLLGNRLFFDQQGDMPMLLDIIQWQW 459
ratT1R2
                HALHSLLGCDKSTCT-KRVVYPWQLLEEIWKVNFTLLDHQIFFDPQGDVALHLEIVQWQW 455
humanT1R2
                                   * ***** :: :: *
                                                         : *: . .
humanT1R3
                QGSVPRLHDVGRFNG---SLRTERLKIRWHTSDNQKPVSRCSRQCQEGQVRRVKGFHSCC 518
catT1R3
                QDPTPELRTVGTFKG---RLELWRSQMCWHTPGKQQPVSQCSRQCKEGQVRRVKGFHSCC 521
                QSPTPVLHTVGTFNG---TLQLQQSKMYWP--GNQVPVSQCSRQCKDGQVRRVKGFHSCC 523
mouseT1R3
ratT1R3
                QSPTPVLHTVGTFNG---TLQLQHSKMYWP--GNQVPVSQCSRQCKDGOVRRVKGFHSCC 523
                NGPEWTFEVIGSASLSPVHLDINKTKIQWHGKNNQVPVSVCTRDCLEGHHRLVMGSHHCC 517
mouseT1R1
                NGPEWTFEIIGSASLSPVHLDINKTKIQWHGKNNQVPVSVCTTDCLAGHHRVVVGSHHCC 515
ratT1R1
humanT1R1
                NGPKWTFTVLGSSTWSPVQLNINETKIQWHGKDNQVPKSVCSSDCLEGHQRVVTGFHHCC 516
mouseT1R2
                GLSQNPFQSIASYSPTETRLTY-ISNVSWYTPNNTVPISMCSKSCQPGQMKKPIGLHPCC 518
ratT1R2
                DLSQNPFQSIASYSPTSKRLTY-INNVSWYTPNNTVPVSMCSKSCQPGQMKKSVGLHPCC 518
humanT1R2
                DRSQNPFQSVASYYPLQRQLKN-IQDISWHTVNNTIPMSMCSKRCQSGQKKKPVGIHVCC 514
                                          .: *
                                                . :
humanT1R3
                YDCVDCEAGSYRQ-NPDDIACTFCGQDEWSPERSTRCFRRRSRFLAWGEPAVLLLLLLS 577
                YNCVDCKAGSYQR-NPDDLLCTQCDQDQWSPDRSTRCFARKPMFLAWGEPAVLLLLALLA 580
catT1R3
mouseT1R3
                YDCVDCKAGSYRK-HPDDFTCTPCNQDQWSPEKSTACLPRRPKFLAWGEPVVLSLLLLLC 582
                YDCVDCKAGSYRK-HPDDFTCTPCGKDQWSPEKSTTCLPRRPKFLAWGEPAVLSLLLLLC 582
ratT1R3
mouseT1R1
                FECMPCEAGTFLN-TSELHTCQPCGTEEWAPEGSSACFSRTVEFLGWHEPISLVLLAANT 576
ratTlRl
                FECVPCEAGTFLN-MSELHICQPCGTEEWAPKESTTCFPRTVEFLAWHEPISLVLIAANT 574
humanT1R1
                FECVPCGAGTFLN-KSDLYRCQPCGKEEWAPEGSQTCFPRTVVFLALREHTSWVLLAANT 575
                FECVDCPPGTYLNRSVDEFNCLSCPGSMWSYKNNIACFKRRLAFLEWHEVPTIVVTILAA 578
mouseT1R2
ratT1R2
                FECLDCMPGTYLNRSADEFNCLSCPGSMWSYKNDITCFQRRPTFLEWHEVPTIVVAILAA 578
                FECIDCLPGTFLNHTEDEYECQACPNNEWSYQSETSCFKRQLVFLEWHEAPTIAVALLAA 574
humanT1R2
                                          . *: . .
                                                    *: *
humanT1R3
                LALGLVLAALGLFVHHRDSPLVQASGGPLACFGLVCLGLVCLSVLLFPGQPSPARCLAQQ 637
catT1R3
                LALGLALAALGLFLWHSDSPLVQASGGPRACFGLACLGLVCLSVLLFPGQPGPASCLAQQ 640
mouseT1R3
                LVLGLALAALGLSVHHWDSPLVQASGGSQFCFGLICLGLFCLSVLLFPGRPSSASCLAQQ 642
ratT1R3
                LVLGLTLAALGLFVHYWDSPLVQASGGSLFCFGLICLGLFCLSVLLFPGRPRSASCLAQQ 642
mouseT1R1
                LLLLLIGTAGLFAWRLHTPVVRSAGGRLCFLMLGSLVAGSCSLYSFFGKPTVPACLLRQ 636
ratT1R1
                LLLLLLVGTAGLFAWHFHTPVVRSAGGRLCFLMLGSLVAGSCSFYSFFGEPTVPACLLRO 634
                LLLLLLLGTAGLFAWHLDTPVVRSAGGRLCFLMLGSLAAGSGSLYGFFGEPTRPACLLRQ 635
humanT1R1
mouseT1R2
                LGFISTLAILLIFWRHFQTPMVRSAGGPMCFLMLVPLLLAFGMVPVYVGPPTVFSCFCRQ 638
ratT1R2
                LGFFSTLAILFIFWRHFQTPMVRSAGGPMCFLMLVPLLLAFGMVPVYVGPPTVFSCFCRQ 638
humanT1R2
                LGFLSTLAILVIFWRHFQTPIVRSAGGPMCFLMLTLLLVAYMVVPVYVGPPKVSTCLCRQ 634
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App No.: Not Yet Assigned

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Title: TASTE RECEPTOR OF THE TIR FAMILY FROM DOMESTIC CAT
Inventors: Xia LI, Weihua LI, Danielle R. REED, Alexander Alexeyevich BACHMANOV,
Joseph G. BRAND

Attorney: Felicity E. Groth
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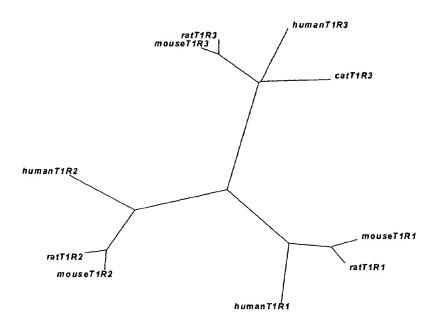
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## Figure 2C

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humanT1R1 mouseT1R2 ratT1R2 humanT1R2	ILCRPDLNSTEHFQASIQDYTRRCGST	

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Figure 3
Phylogenetic Tree of T1Rs:



0.1

App No.: Not Yet Assigned Filed: Herewith Title: TASTE RECEPTOR OF THE TIR FAMILY FROM DOMESTIC CAT Inventors: Xia LI, Weihua LI, Danielle R. REED, Alexander Alexeyevich BACHMANOV, Joseph G. BRAND

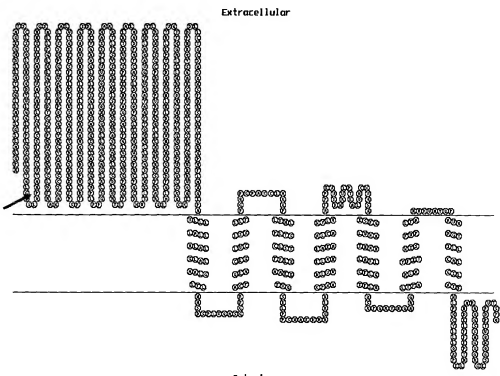
Attorney: Felicity E. Groth Sheet 13 of 13

Phone: (215) 568-3100

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Figure 4. Predicted conformation of the 7TM T1R3 protein sequence from cat.

Arrow points to region of possible functional amino acide substitution.



Cytoplasm

## SEQUENCE LISTING

<110> Li, Xia Li, Weihua Reed, Danielle Bachmanov, Alexander Brand, Joseph	Li, Weihua Reed, Danielle Bachmanov, Alexander												
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Ala Gly Leu Arg Ser Arg Thr Arg Pro Ser Ser Pro Val Cys Thr Arg 50 60

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Page 19

695 700

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Cys	Asn	Val	Ser 420	His	Cys	His	Thr	Ser 425	Glu	Pro	Val	Gln	Pro 430	Trp	Gln
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Leu	Gln 450	Phe	Asp	Ala	Lys	Gly 455	Ser	Val	Asp	Met	Glu 460	Tyr	Asp	Leu	Lys
Met 465	Trp	Val	Trp	Gln	Ser 470	Pro	Thr	Pro	Val	Leu 475	His	Thr	Val	Gly	Thr 480
Phe	Asn	Gly	Thr	Leu 485	Gln	Leu	Gln	His	Ser 490	Lys	Met	Туr	Trp	Pro 495	Gly
Asn	Gln	Val	Pro 500	Val	Ser	Gln	Cys	Ser 505	Arg	Gln	Cys	Lys	Asp 510	Gly	Gln
Val	Arg	Arg 515	Val	Lys	Gly	Phe	His 520	Ser	Cys	Суз	Tyr	Asp 525	Cys	Val	Asp
Cys	<b>Lys</b> 530	Ala	Gly	Ser	Tyr	Arg 535	Lys	His	Pro	Asp	Asp 540	Phe	Thr	Cys	Thr
Pro 545	Cys	Gly	Lys	Asp	Gln 550	Trp	Ser	Pro	Glu	Lys 555	Ser	Thr	Thr	Cys	Leu 560
Pro	Arg	Arg	Pro	Lys 565	Phe	Leu	Ala	Trp	Gly 570	Glu	Pro	Ala	Val	Leu 575	Ser
Leu	Leu	Leu	Leu 580	Leu	Cys	Leu	Val	Leu 585	Gly	Leu	Thr	Leu	Ala 590	Ala	Leu
Gly	Leu	Phe 595	Val	His	Tyr	Trp	Asp 600	Ser	Pro	Leu	Val	Gln 605	Ala	Ser	Gly
Gly	Ser	Leu	Phe	Cys	Phe	Gly	Leu	Ile	Cys		Gly ge 2	_	Phe	Cys	Leu

.

610 615 620

Ser Val Leu Leu Phe Pro Gly Arg Pro Arg Ser Ala Ser Cys Leu Ala

Gln Gln Pro Met Ala His Leu Pro Leu Thr Gly Cys Leu Ser Thr Leu

Phe Leu Gln Ala Ala Glu Ile Phe Val Glu Ser Glu Leu Pro Leu Ser

Trp Ala Asn Trp Leu Cys Ser Tyr Leu Arg Gly Pro Trp Ala Trp Leu

Val Val Leu Leu Ala Thr Leu Val Glu Ala Ala Leu Cys Ala Trp Tyr

Leu Met Ala Phe Pro Pro Glu Val Val Thr Asp Trp Gln Val Leu Pro 705 710 715 720

Thr Glu Val Leu Glu His Cys Arg Met Arg Ser Trp Val Ser Leu Gly  $725 \hspace{1cm} 730 \hspace{1cm} 735$ 

Leu Val His Ile Thr Asn Ala Val Leu Ala Phe Leu Cys Phe Leu Gly

Thr Phe Leu Val Gln Ser Gln Pro Gly Arg Tyr Asn Arg Ala Arg Gly

Leu Thr Phe Ala Met Leu Ala Tyr Phe Ile Ile Trp Val Ser Phe Val 770 780

Pro Leu Leu Ala Asn Val Gln Val Ala Tyr Gln Pro Ala Val Gln Met

Gly Ala Ile Leu Phe Cys Ala Leu Gly Ile Leu Ala Thr Phe His Leu 810

Pro Lys Cys Tyr Val Leu Leu Trp Leu Pro Glu Leu Asn Thr Gln Glu 820 825 830

Phe Phe Leu Gly Arg Ser Pro Lys Glu Ala Ser Asp Gly Asn Ser Gly

Ser Ser Glu Ala Thr Arg Gly His Ser Glu

<210> 15 <211> 842

<212> PRT

<213> Mus musculus

<400> 15

Met Leu Phe Trp Ala Ala His Leu Leu Ser Leu Gln Leu Ala Val

Ala Tyr Cys Trp Ala Phe Ser Cys Gln Arg Thr Glu Ser Ser Pro Gly 20 25 30

Phe Ser Leu Pro Gly Asp Phe Leu Leu Ala Gly Leu Phe Ser Leu His  $35 \hspace{1cm} 40 \hspace{1cm} 45$ 

Ala Asp Cys Leu Gln Val Arg His Arg Pro Leu Val Thr Ser Cys Asp 50 60

Arg Ser Asp Ser Phe Asn Gly His Gly Tyr His Leu Phe Gln Ala Met 70 75 80

Arg Phe Thr Val Glu Glu Ile Asn Asn Ser Thr Ala Leu Leu Pro Asn 85 90 95

Ile Thr Leu Gly Tyr Glu Leu Tyr Asp Val Cys Ser Glu Ser Ser Asn 100 105 110

Val Tyr Ala Thr Leu Arg Val Leu Ala Gln Gln Gly Thr Gly His Leu 115 120 125

Glu Met Gln Arg Asp Leu Arg Asn His Ser Ser Lys Val Val Ala Leu 130 135 140

Ile Gly Pro Asp Asn Thr Asp His Ala Val Thr Thr Ala Ala Leu Leu 145 150 160

Ser Pro Phe Leu Met Pro Leu Val Ser Tyr Glu Ala Ser Ser Val Ile 165 170 175

Leu Ser Gly Lys Arg Lys Phe Pro Ser Phe Leu Arg Thr Ile Pro Ser 180 185 190

Asp Lys Tyr Gln Val Glu Val Ile Val Arg Leu Leu Gln Ser Phe Gly 195 200 205

Trp Val Trp Ile Ser Leu Val Gly Ser Tyr Gly Asp Tyr Gly Gln Leu 210 220

Gly Val Gln Ala Leu Glu Glu Leu Ala Thr Pro Arg Gly Ile Cys Val 225 230 235 240

Ala Phe Lys Asp Val Val Pro Leu Ser Ala Gln Ala Gly Asp Pro Arg 245 250 255

Met Gln Arg Met Met Leu Arg Leu Ala Arg Ala Arg Thr Thr Val Val 260 265 270

Val Val Phe Ser Asn Arg His Leu Ala Gly Val Phe Phe Arg Ser Val 275 280 285

Val	Leu 290	Ala	Asn	Leu	Thr	Gly 295	Lys	Val	Trp	Ile	Ala 300	Ser	Glu	Asp	Trp
Ala 305	Ile	Ser	Thr	Tyr	11e 310	Thr	Asn	Val	Pro	Gly 315	Ile	Gln	Gly	Ile	Gly 320
Thr	Val	Leu	Gly	Val 325	Ala	Ile	Gln	Gln	Arg 330	Gln	Val	Pro	Gly	Leu 335	Lys
Glu	Phe	Glu	Glu 340	Ser	Tyr	Val	Gln	Ala 345	Val	Met	Gly	Ala	Pro 350	Arg	Thr
Cys	Pro	Glu 355	Gly	Ser	Trp	Cys	Gly 360	Thr	Asn	Gln	Leu	Cys 365	Arg	Glu	Cys
His	Ala 370	Phe	Thr	Thr	Trp	Asn 375	Met	Pro	Glu	Leu	Gly 380	Ala	Phe	Ser	Met
Ser 385	Ala	Ala	Tyr	Asn	Val 390	Tyr	Glu	Ala	Val	Tyr 395	Ala	Val	Ala	His	Gly 400
Leu	His	Gln	Leu	Leu 405	Gly	Cys	Thr	Ser	Gly 410	Thr	Суѕ	Ala	Arg	Gly 415	Pro
Val	Tyr	Pro	Trp 420	Gln	Leu	Leu	Gln	Gln 425	Ile	Tyr	Lys	Val	Asn 430	Phe	Leu
Leu	His	Lys 435	Lys	Thr	Val	Ala	Phe 440	Asp	Asp	Lys	Gly	Asp 445	Pro	Leu	Gly
Tyr	Tyr 450	Asp	Ile	Ile	Ala	Trp 455	Asp	Trp	Asn	Gly	Pro 460	Glu	Trp	Thr	Phe
Glu 465	Val	Ile	Gly	Ser	Ala 470	Ser	Leu	Ser	Pro	Val 475	His	Leu	Asp	Ile	Asn 480
Lys	Thr	Lys	Ile	Gln 485	Trp	His	Gly	Lys	Asn 490	Asn	Gln	Val	Pro	Val 495	Ser
Val	Cys	Thr	Arg 500	Asp	Cys	Leu	Glu	Gly 505	His	His	Arg	Leu	Val 510	Met	Gly
Ser	His	His 515	Cys	Cys	Phe	<b>Gl</b> u	Cys 520	Met	Pro	Суз	Glu	Ala 525	Gly	Thr	Phe
Leu	Asn 530	Thr	Ser	Glu	Leu	His 535	Thr	Cys	Gln	Pro	Cys 540	Gly	Thr	Glu	Glu
Trp 545	Ala	Pro	Glu	Gly	Ser 550	Ser	Ala	Cys	Phe	Ser 555	Arg	Thr	Val	Glu	Phe 560
Leu	Gly	Trp	His	Glu 565	Pro	Ile	Ser	Leu	Val 570	Leu	Leu	Ala	Ala	Asn 575	Thr

Leu Leu Leu Leu Leu Ile Gly Thr Ala Gly Leu Phe Ala Trp Arg Leu His Thr Fro Val Val Arg Ser Ala Gly Gly Arg Leu Cys Phe Leu Met Leu Gly Ser Leu Val Ala Gly Ser Cys Ser Leu Tyr Ser Phe Phe Gly Lys Pro Thr Val Pro Ala Cys Leu Leu Arg Gln Pro Leu Phe Ser Leu Gly Phe Ala Ile Phe Leu Ser Cys Leu Thr Ile Arg Ser Phe Gln Leu Val Ile Ile Phe Lys Phe Ser Thr Lys Val Pro Thr Phe Tyr His Thr Trp Ala Gln Asn His Gly Ala Gly Ile Phe Val Ile Val Ser Ser Thr Val His Leu Phe Leu Cys Leu Thr Trp Leu Ala Met Trp Thr Pro Arg Pro Thr Arg Glu Tyr Gln Arg Phe Pro His Leu Val Ile Leu Glu Cys Thr Glu Val Asn Ser Val Gly Phe Leu Val Ala Phe Ala His Asn Ile Leu Leu Ser Ile Ser Thr Phe Val Cys Ser Tyr Leu Gly Lys Glu 740 745 750Leu Pro Glu Asn Tyr Asn Glu Ala Lys Cys Val Thr Phe Ser Leu Leu Leu His Phe Val Ser Trp Ile Ala Phe Phe Thr Met Ser Ser Ile Tyr 770 780Gln Gly Ser Tyr Leu Pro Ala Val Asn Val Leu Ala Gly Leu Ala Thr Leu Ser Gly Gly Phe Ser Gly Tyr Phe Leu Pro Lys Cys Tyr Val Ile Leu Cys Arg Pro Glu Leu Asn Asn Thr Glu His Phe Gln Ala Ser Ile 825 Gln Asp Tyr Thr Arg Arg Cys Gly Thr Thr

<210> 16 <211> 840

<212> PRT <213> Rattus rattus

<400> 16

Met Leu Phe Trp Ala Ala His Leu Leu Leu Ser Leu Gln Leu Val Tyr

Cys Trp Ala Phe Ser Cys Gln Arg Thr Glu Ser Ser Pro Gly Phe Ser 20 25 30

Leu Pro Gly Asp Phe Leu Leu Ala Gly Leu Phe Ser Leu His Gly Asp

Cys Leu Gln Val Arg His Arg Pro Leu Val Thr Ser Cys Asp Arg Pro

Asp Ser Phe Asn Gly His Gly Tyr His Leu Phe Gln Ala Met Arg Phe

Thr Val Glu Glu Ile Asn Asn Ser Ser Ala Leu Leu Pro Asn Ile Thr

Leu Gly Tyr Glu Leu Tyr Asp Val Cys Ser Glu Ser Ala Asn Val Tyr

Ala Thr Leu Arg Val Leu Ala Leu Gln Gly Pro Arg His Ile Glu Ile

Gln Lys Asp Leu Arg Asn His Ser Ser Lys Val Val Ala Phe Ile Gly 135

Pro Asp Asn Thr Asp His Ala Val Thr Thr Ala Ala Leu Leu Gly Pro

Phe Leu Met Pro Leu Val Ser Tyr Glu Ala Ser Ser Val Val Leu Ser

Ala Lys Arg Lys Phe Pro Ser Phe Leu Arg Thr Val Pro Ser Asp Arg

His Gln Val Glu Val Met Val Gln Leu Leu Gln Ser Phe Gly Trp Val 200

Trp Ile Ser Leu Ile Gly Ser Tyr Gly Asp Tyr Gly Gln Leu Gly Val

Gln Ala Leu Glu Glu Leu Ala Val Pro Arg Gly Ile Cys Val Ala Phe

Lys Asp Ile Val Pro Phe Ser Ala Arg Val Gly Asp Pro Arg Met Gln

Ser Met Met Gln His Leu Ala Gln Ala Arg Thr Thr Val Val Val Val 265

Phe Ser Asn Arg His Leu Ala Arg Val Phe Phe Arg Ser Val Val Leu Ala Asn Leu Thr Gly Lys Val Trp Val Ala Ser Glu Asp Trp Ala Ile Ser Thr Tyr Ile Thr Ser Val Thr Gly Ile Gln Gly Ile Gly Thr Val Leu Gly Val Ala Val Gln Gln Arg Gln Val Pro Gly Leu Lys Glu Phe Glu Glu Ser Tyr Val Arg Ala Val Thr Ala Ala Pro Ser Ala Cys Pro Glu Gly Ser Trp Cys Ser Thr Asn Gln Leu Cys Arg Glu Cys His Thr Phe Thr Thr Arg Asn Met Pro Thr Leu Gly Ala Phe Ser Met Ser Ala Ala Tyr Arg Val Tyr Glu Ala Val Tyr Ala Val Ala His Gly Leu His Gln Leu Leu Gly Cys Thr Ser Glu Ile Cys Ser Arg Gly Pro Val Tyr Pro Trp Gln Leu Leu Gln Gln Ile Tyr Lys Val Asn Phe Leu Leu His 420 425 430Glu Asn Thr Val Ala Phe Asp Asp Asn Gly Asp Thr Leu Gly Tyr Tyr Asp Ile Ile Ala Trp Asp Trp Asn Gly Pro Glu Trp Thr Phe Glu Ile Ile Gly Ser Ala Ser Leu Ser Pro Val His Leu Asp Ile Asn Lys Thr Lys Ile Gln Trp His Gly Lys Asn Asn Gln Val Pro Val Ser Val Cys Thr Thr Asp Cys Leu Ala Gly His His Arg Val Val Val Gly Ser His His Cys Cys Phe Glu Cys Val Pro Cys Glu Ala Gly Thr Phe Leu Asn Met Ser Glu Leu His Ile Cys Gln Pro Cys Gly Thr Glu Glu Trp Ala 530 535 Pro Lys Glu Ser Thr Thr Cys Phe Pro Arg Thr Val Glu Phe Leu Ala Page 31

Trp His Glu Pro Ile Ser Leu Val Leu Ile Ala Ala Asn Thr Leu Leu 565 570 575

Leu Leu Leu Val Gly Thr Ala Gly Leu Phe Ala Trp His Phe His 580 585 590

Thr Pro Val Val Arg Ser Ala Gly Gly Arg Leu Cys Phe Leu Met Leu 595 600 605

Gly Ser Leu Val Ala Gly Ser Cys Ser Phe Tyr Ser Phe Phe Gly Glu 610 615 620

Pro Thr Val Pro Ala Cys Leu Leu Arg Gln Pro Leu Phe Ser Leu Gly 625 630 635 640

Phe Ala Ile Phe Leu Ser Cys Leu Thr Ile Arg Ser Phe Gln Leu Val 645 650 655

Ile Ile Phe Lys Phe Ser Thr Lys Val Pro Thr Phe Tyr Arg Thr Trp
660 665 670

Ala Gln Asn His Gly Ala Gly Leu Phe Val Ile Val Ser Ser Thr Val 675 680 685

His Leu Leu Ile Cys Leu Thr Trp Leu Val Met Trp Thr Pro Arg Pro 690 700

Thr Arg Glu Tyr Gln Arg Phe Pro His Leu Val Ile Leu Glu Cys Thr 705 710 715 720

Glu Val Asn Ser Val Gly Phe Leu Leu Ala Phe Thr His Asn Ile Leu 725 730 735

Glu Asn Tyr Asn Glu Ala Lys Cys Val Thr Phe Ser Leu Leu Leu Asn 755 760 765

Phe Val Ser Trp Ile Ala Phe Phe Thr Met Ala Ser Ile Tyr Gln Gly 770 780

Ser Tyr Leu Pro Ala Val Asn Val Leu Ala Gly Leu Thr Thr Leu Ser 785 790 795 800

Gly Gly Phe Ser Gly Tyr Phe Leu Pro Lys Cys Tyr Val Ile Leu Cys 805 810 815

Arg Pro Glu Leu Asn Asn Thr Glu His Phe Gln Ala Ser Ile Gln Asp 820 825 830

Tyr Thr Arg Arg Cys Gly Thr Thr <210> 17 <211> 841 <212> PRT <213> Homo sapiens <400> 17 Met Leu Leu Cys Thr Ala Arg Leu Val Gly Leu Gln Leu Leu Ile Ser Cys Cys Trp Ala Phe Ala Cys His Ser Thr Glu Ser Ser Pro Asp Phe Thr Leu Pro Gly Asp Tyr Leu Leu Ala Gly Leu Phe Pro Leu His Ser Gly Cys Leu Gln Val Arg His Arg Pro Glu Val Thr Leu Cys Asp Arg 50 60 Ser Cys Ser Phe Asn Glu His Gly Tyr His Leu Phe Gln Ala Met Arg Leu Gly Val Glu Glu Ile Asn Asn Ser Thr Ala Leu Leu Pro Asn Ile Thr Leu Gly Tyr Gln Leu Tyr Asp Val Cys Ser Asp Ser Ala Asn Val Tyr Ala Thr Leu Arg Val Leu Ser Leu Pro Gly Gln His His Ile Glu Leu Gln Gly Asp Leu Leu His Tyr Ser Pro Thr Val Leu Ala Val Ile Gly Pro Asp Ser Thr Asn Arg Ala Ala Thr Thr Ala Ala Leu Leu Ser Pro Phe Leu Val Pro Met Ile Ser Tyr Ala Ala Ser Ser Glu Thr Leu Ser Val Lys Arg Gln Tyr Pro Ser Phe Leu Arg Thr Ile Pro Asn Asp Lys Tyr Gln Val Glu Thr Met Val Leu Leu Gln Lys Phe Gly Trp 200 Thr Trp Ile Ser Leu Val Gly Ser Ser Asp Asp Tyr Gly Gln Leu Gly

Val Gln Ala Leu Glu Asn Gln Ala Thr Gly Gln Gly Ile Cys Ile Ala

Phe	Lys	Asp	Ile	Met 245	Pro	Phe	Ser	Ala	Gln 250	Val	Gly	Asp	Glu	Arg 255	Met
Gln	Cys	Leu	Met 260	Arg	His	Leu	Ala	Gln 265	Ala	Gly	Ala	Thr	Val 270	Val	Val
Val	Phe	Ser 275	Ser	Arg	Gln	Leu	Ala 280	Arg	Val	Phe	Phe	Glu 285	Ser	Val	Val
Leu	Thr 290	Asn	Leu	Thr	Gly	Lys 295	Val	Trp	Val	Ala	Ser 300	Glu	Ala	Trp	Ala
Leu 305	Ser	Arg	His	Ile	Thr 310	Gly	Val	Pro	Gly	Ile 315	Gln	Arg	Ile	Gly	Met 320
Val	Leu	Gly	Val	Ala 325	Ile	Gln	Lys	Arg	Ala 330	Val	Pro	Gly	Leu	Lys 335	Ala
Phe	Glu	Glu	Ala 340	Tyr	Ala	Arg	Ala	Asp 345	Lys	Lys	Ala	Pro	Arg 350	Pro	Cys
His	Lys	Gly 355	Ser	Trp	Cys	Ser	Ser 360	Asn	Gln	Leu	Cys	Arg 365	Glu	Cys	Gln
Ala	Phe 370	Met	Ala	His	Thr	Met 375	Pro	Lys	Leu	Lys	Ala 380	Phe	Ser	Met	Ser
Ser 385	Ala	Tyr	Asn	Ala	Tyr 390	Arg	Ala	Val	Tyr	Ala 395	Val	Ala	His	Gly	Leu 400
His	Gln	Leu	Leu	Gly 405	Cys	Ala	Ser	Gly	Ala 410	Суѕ	Ser	Arg	Gly	Arg 415	Val
Tyr	Pro	Trp	Gln 420	Leu	Leu	Glu	Gln	Ile 425	His	Lys	Val	His	Phe 430	Leu	Leu
His	Lys	Asp 435	Thr	Val	Ala	Phe	Asn 440	Asp	Asn	Arg	Asp	Pro 445	Leu	Ser	Ser
Tyr	Asn 450	Ile	Ile	Ala	Trp	Asp 455	Trp	Asn	Gly	Pro	Lys 460	Trp	Thr	Phe	Thr
Val 465	Leu	Gly	Ser	Ser	Thr 470	Trp	Ser	Pro	Val	Gln 475	Leu	Asn	Ile	Asn	Glu 480
Thr	Lys	Ile	Gln	Trp 485	His	Gly	Lys	Asp	Asn 490	Gln	Val	Pro	Lys	Ser 495	Val
Cys	Ser	Ser	Asp 500	Cys	Leu	Glu	Gly	His 505	Gln	Arg	Val	Val	Thr 510	Gly	Phe
His	His	Cys 515	Cys	Phe	Glu	Суѕ	Val 520	Pro	Cys	Gly	Ala	Gly 525	Thr	Phe	Leu

Asn	Lys 530	Ser	Asp	Leu	Туr	Arg 535	Cys	Gln	Pro	Cys	Gly 540	Lys	Glu	Glu	Trp
Ala 545	Pro	Glu	Gly	Ser	Gln 550	Thr	Cys	Phe	Pro	Arg 555	Thr	Val	Val	Phe	Leu 560
Ala	Leu	Arg	Glu	His 565	Thr	Ser	Trp	Val	Leu 570	Leu	Ala	Ala	Asn	Thr 575	Leu
Leu	Leu	Leu	Leu 580	Leu	Leu	Gly	Thr	Ala 585	Gly	Leu	Phe	Ala	Trp 590	His	Leu
Asp	Thr	Pro 595	Val	Val	Arg	Ser	Ala 600	Gly	Gly	Arg	Leu	Cys 605	Phe	Leu	Met
Leu	Gly 610	Ser	Leu	Ala	Ala	Gly 615	Ser	Gly	Ser	Leu	Tyr 620	Gly	Phe	Phe	Gly
Glu 625	Pro	Thr	Arg	Pro	Ala 630	Cys	Leu	Leu	Arg	Gln 635	Ala	Leu	Phe	Ala	Leu 640
Gly	Phe	Thr	Ile	Phe 645	Leu	Ser	Суѕ	Leu	Thr 650	Val	Arg	Ser	Phe	Gln 655	Leu
Ile	Ile	Ile	Phe 660	Lys	Phe	Ser	Thr	Lys 665	Val	Pro	Thr	Phe	Tyr 670	His	Ala
Trp	Val	Gln 675	Asn	His	Gly	Ala	Gly 680	Leu	Phe	Val	Met	Ile 685	Ser	Ser	Ala
Ala	Gln 690	Leu	Leu	Ile	Cys	Leu 695	Thr	Trp	Leu	Val	Val 700	Trp	Thr	Pro	Leu
Pro 705	Ala	Arg	Glu	Tyr	Gln 710	Arg	Phe	Pro	His	Leu 715	Val	Met	Leu	Glu	Cys 720
Thr	Glu	Thr	Asn	Ser 725	Leu	Gly	Phe	Ile	Leu 730	Ala	Phe	Leu	Tyr	Asn 735	Gly
Leu	Leu	Ser	Ile 740	Ser	Ala	Phe	Ala	Cys 745	Ser	Tyr	Leu	Gly	Lys 750	Asp	Leu
Pro	Glu	Asn 755	Tyr	Asn	Glu	Ala	Lys 760	Суѕ	Val	Thr	Phe	Ser 765	Leu	Leu	Phe
Asn	Phe 770	Val	Ser	Trp	Ile	Ala 775	Phe	Phe	Thr	Thr	Ala 780	Ser	Val	Tyr	Asp
Gly 785	Lys	Tyr	Leu	Pro	Ala 790	Ala	Asn	Met	Met	Ala 795	Gly	Leu	Ser	Ser	Leu 800
Ser	Ser	Gly	Phe	Gly 805	Gly	Tyr	Phe	Leu	Pro 810	Lys	Cys	Tyr	Val	Ile 815	Leu

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Ile Val Val Leu Val Ser Asp Asp Asp Tyr Gly Arg Glu Asn Ser His 210 215 220

His Ile Glu Ala Met Val Gln Leu Met Val His Phe Gln Trp Asn Trp

Leu 225	Leu	Ser	Gln	Arg	Leu 230	Thr	Asn	Thr	Gly	Asp 235	Ile	Cys	Ile	Ala	Phe 240
Gln	Glu	Val	Leu	Pro 245	Val	Pro	Glu	Pro	Asn 250	Gln	Ala	Val	Arg	Pro 255	Glu
Glu	Gln	Asp	Gln 260	Leu	Asp	Asn	Ile	Leu 265	Asp	Lys	Leu	Arg	Arg 270	Thr	Ser
Ala	Arg	Val 275	Val	Val	Ile	Phe	Ser 280	Pro	Glu	Leu	Ser	Leu 285	His	Asn	Phe
Phe	Arg 290	Glu	Val	Leu	Arg	Trp 295	Asn	Phe	Thr	Gly	Phe 300	Val	Trp	Ile	Ala
Ser 305	Glu	Ser	Trp	Ala	Ile 310	Asp	Pro	Val	Leu	His 315	Asn	Leu	Thr	Glu	Leu 320
Arg	His	Thr	Gly	Thr 325	Phe	Leu	Gly	Val	Thr 330	Ile	Gln	Arg	Val	Ser 335	Ile
Pro	Gly	Phe	Ser 340	Gln	Phe	Arg	Val	Arg 345	His	Asp	Lys	Pro	Glu 350	Туг	Pro
Met	Pro	Asn 355	Glu	Thr	Ser		Arg 360	Thr	Thr	Суѕ	Asn	Gln 365	Asp	Cys	Asp
Ala	Cys 370	Met	Asn	Ile	Thr	Glu 375	Ser	Phe	Asn	Asn	Val 380	Leu	Met	Leu	Ser
Gly 385	Glu	Arg	Val	Val	Туг 390	Ser	Val	Tyr	Ser	Ala 395	Val	Tyr	Ala	Val	Ala 400
His	Thr	Leu	His	Arg 405	Leu	Leu	His	Cys	Asn 410	Gln	Val	Arg	Cys	Thr 415	Lys
Gln	Ile	Val	Tyr 420	Pro	Trp	Gln	Leu	Leu 425	Arg	Glu	Ile	Trp	His 430	Val	Asn
Phe	Thr	Leu 435	Leu	Gly	Asn	Gln	Leu 440	Phe	Phe	Asp	Glu	Gln 445	Gly	Asp	Met
Pro	Met 450	Leu	Leu	Asp	Ile	Ile 455	Gln	Trp	Gln	Trp	Gly 460	Leu	Ser	Gln	Asn
Pro 465	Phe	Gln	Ser	Ile	Ala 470	Ser	Tyr	Ser	Pro	Thr 475	Glu	Thr	Arg	Leu	Thr 480
Tyr	Ile	Ser	Asn	Val 485	Ser	Trp	Tyr	Thr	Pro 490	Asn	Asn	Thr	Val	Pro 495	Ile
Ser	Met	Cys	Ser	Lys	Ser	Суѕ	Gln	Pro	Gly		Met ge 3	_	Lys	Pro	Ile

Gly Leu His Pro Cys Cys Phe Glu Cys Val Asp Cys Pro Pro Gly Thr 515 520 525

Tyr Leu Asn Arg Ser Val Asp Glu Phe Asn Cys Leu Ser Cys Pro Gly 530 540

Ser Met Trp Ser Tyr Lys Asn Asn Ile Ala Cys Phe Lys Arg Arg Leu 545 550 555 560

Ala Phe Leu Glu Trp His Glu Val Pro Thr Ile Val Val Thr Ile Leu 565 570 575

Ala Ala Leu Gly Phe Ile Ser Thr Leu Ala Ile Leu Leu Ile Phe Trp 580 585 590

Arg His Phe Gln Thr Pro Met Val Arg Ser Ala Gly Gly Pro Met Cys 595 600 605

Phe Leu Met Leu Val Pro Leu Leu Leu Ala Phe Gly Met Val Pro Val 610 620

Tyr Val Gly Pro Pro Thr Val Phe Ser Cys Phe Cys Arg Gln Ala Phe 625 630 635

Phe Thr Val Cys Phe Ser Val Cys Leu Ser Cys Ile Thr Val Arg Ser 645 650 655

Phe Gln Ile Val Cys Val Phe Lys Met Ala Arg Arg Leu Pro Ser Ala 660 665 670

Tyr Gly Phe Trp Met Arg Tyr His Gly Pro Tyr Val Phe Val Ala Phe 675 680 685

Ile Thr Ala Val Lys Val Ala Leu Val Ala Gly Asn Met Leu Ala Thr 690 695 700

Thr Ile Asn Pro Ile Gly Arg Thr Asp Pro Asp Asp Pro Asn Ile Ile 705 710 715 720

Ile Leu Ser Cys His Pro Asn Tyr Arg Asn Gly Leu Leu Phe Asn Thr

Ser Met Asp Leu Leu Ser Val Leu Gly Phe Ser Phe Ala Tyr Val
740 745 750

Gly Lys Glu Leu Pro Thr Asn Tyr Asn Glu Ala Lys Phe Ile Thr Leu 755 760 765

Ser Met Thr Phe Ser Phe Thr Ser Ser Ile Ser Leu Cys Thr Phe Met

Ser Val His 785	Asp Gly	Val Leu 790	Val	Thr	Ile	Met 795	Asp	Leu	Leu	Val	Thr 800
Val Leu Asn	Phe Leu 805	Ala Ile	Gly	Leu	Gly 810	Tyr	Phe	Gly	Pro	Lys 815	Cys
Tyr Met Ile	Leu Phe 820	Tyr Pro	Glu	Arg 825	Asn	Thr	Ser	Ala	Tyr 830	Phe	Asn
Ser Met Ile 835	Gln Gly	Tyr Thr	Met 840	Arg	Lys	Ser					
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Val Leu Pro	Lys Pro 20	Gly Lys	Leu	Val 25	Glu	Asn	Ser	Asp	Phe 30	His	Leu
Ala Gly Asp 35	Tyr Leu	Leu Gly	Gly 40	Leu	Phe	Thr	Leu	His 45	Ala	Asn	Val
Lys Ser Ile 50	Ser His	Leu Ser 55	Tyr	Leu	Gln	Val	Pro 60	Lys	Cys	Asn	Glu
Phe Thr Met 65	Lys Val	Leu Gly 70	Tyr	Asn	Leu	Met 75	Gln	Ala	Met	Arg	Phe 80
Ala Val Glu	Glu Ile 85	Asn Asn	Cys	Ser	Ser 90	Leu	Leu	Pro	Gly	Val 95	Leu
Leu Gly Tyr	Glu Met 100	Val Asp	Val	Cys 105	Tyr	Leu	Ser	Asn	Asn 110	Ile	His
Pro Gly Leu 115	Tyr Phe	Leu Ala	Gln 120	Asp	Asp	Asp	Leu	Leu 125	Pro	Ile	Leu
Lys Asp Tyr 130	Ser Gln	Tyr Met 135	Pro	His	Val	Val	Ala 140	Val	Ile	Gly	Pro
Asp Asn Ser 145	Glu Ser	Ala Ile 150	Thr	Val	Ser	Asn 155	Ile	Leu	Ser	His	Phe 160
Leu Ile Pro	Gln Ile 165	Thr Tyr	Ser	Ala	Ile 170	Ser	Asp	Lys	Leu	Arg 175	Asp
Lys Arg His	Phe Pro 180	Ser Met	Leu	Arg 185	Thr	Val	Pro	Ser	Ala 190	Thr	His

His Ile	Glu Ala 195	a Met V	ıl Glm	Leu 200	Met	Val	His	Phe	Gln 205	Trp	Asn	Trp
Ile Val 210	Val Le	ı Val Se	r Asp 215		Asp	Tyr	Gly	Arg 220	Glu	Asn	Ser	His
Leu Leu 225	Ser Gl	a Arg Le 2:		Lys	Thr	Ser	Asp 235	Ile	Cys	Ile	Ala	Phe 240
Gln Glu	Val Le	1 Pro II 245	e Pro	Glu	Ser	Ser 250	Gln	Val	Met	Arg	Ser 255	Glu
Glu Gln	Arg Gla 26		p Asn	Ile	Leu 265	Asp	Lys	Leu	Arg	Arg 270	Thr	Ser
Ala Arg	Val Val 275	l Val Va	l Phe	Ser 280	Pro	Glu	Leu	Ser	Leu 285	Tyr	Ser	Phe
Phe His 290	Glu Va	l Leu A	g Trp 295		Phe	Thr	Gly	Phe 300	Val	Trp	Ile	Ala
Ser Glu 305	Ser Tr	Ala II	_	Pro	Val	Leu	His 315	Asn	Leu	Thr	Glu	Leu 320
Arg His	Thr Gl	y Thr Pl 325	e Leu	Gly	Val	Thr 330	Ile	Gln	Arg	Val	Ser 335	Ile
Pro Gly	Phe Se:		e Arg	Val	Arg 345	Arg	Asp	Lys	Pro	Gly 350	Tyr	Pro
Val Pro	Asn Th: 355	Thr As	n Leu	Arg 360	Thr	Thr	Cys	Asn	Gln 365	Asp	Cys	Asp
Ala Cys 370	Leu Ası	n Thr Th	r Lys 375		Phe	Asn	Asn	11e 380	Leu	Ile	Leu	Ser
Gly Glu 385		3.9	0				395		_			400
His Ala		405			_	410				_	415	_
Gln Lys	420	)			425					430		
Phe Thr	435	_		440					445	_		
Pro Met 450	Leu Lei	ı Asp Il	e Ile 455	Gln	Trp	Gln	Trp	Asp 460	Leu	Ser	Gln	Asn
Pro Phe 465	Gln Se	Ile Al		Tyr	Ser	Pro	Thr 475	Ser	Lys	Arg	Leu	Thr 480

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Tyr	Ile	Asn	Asn	Val 485	Ser	Trp	Tyr	Thr	Pro 490	Asn	Asn	Thr	Val	Pro 495	Val
Ser	Met	Суз	Ser 500	Lys	Ser	Суз	Gln	Pro 505	Gly	Gln	Met	Lys	Lys 510	Ser	Val
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Thr	Phe	Leu	Glu	Trp 565	His	Glu	Val	Pro	Thr 570	Ile	Val	Val	Ala	Ile 575	Leu
Ala	Ala	Leu	Gly 580	Phe	Phe	Ser	Thr	Leu 585	Ala	Ile	Leu	Phe	Ile 590	Phe	Trp
Arg	His	Phe 595	Gln	Thr	Pro	Met	Val 600	Arg	Ser	Ala	Gly	Gly 605	Pro	Met	Cys
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Tyr 625	Val	Gly	Pro	Pro	Thr 630	Val	Phe	Ser	Cys	Phe 635	Cys	Arg	Gln	Ala	Phe 640
Phe	Thr	Val	Cys	Phe 645	Ser	Ile	Суз	Leu	Ser 650	Cys	Ile	Thr	Val	Arg 655	Ser
Phe	Gln	Ile	Val 660	Суз	Val	Phe	Lys	Met 665	Ala	Arg	Arg	Leu	Pro 670	Ser	Ala
Tyr	Ser	Phe 675	Trp	Met	Arg	Tyr	His 680	Gly	Pro	Tyr	Val	Phe 685	Val	Ala	Phe
Ile	Thr 690	Ala	Ile	Lys	Val	Ala 695	Leu	Val	Val	Gly	Asn 700	Met	Leu	Ala	Thr
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Ser	Met	Asp	Leu 740	Leu	Leu	Ser	Val	Leu 745	Gly	Phe	Ser	Phe	Ala 750	Tyr	Met
Gly	Lys	Glu 755	Leu	Pro	Thr	Asn	Tyr 760	Asn	Glu	Ala	Lys	Phe 765	Ile	Thr	Leu

Ser Met Thr Phe Ser Phe Thr Ser Ser Ile Ser Leu Cys Thr Phe Met Ser Val His Asp Gly Val Leu Val Thr Ile Met Asp Leu Leu Val Thr Val Leu Asn Phe Leu Ala Ile Gly Leu Gly Tyr Phe Gly Pro Lys Cys Tyr Met Ile Leu Phe Tyr Pro Glu Arg Asn Thr Ser Ala Tyr Phe Asn Ser Met Ile Gln Gly Tyr Thr Met Arg Lys Ser <210> 20 <211> 839 <212> PRT <213> Homo sapiens <400> 20 Met Gly Pro Arg Ala Lys Thr Ile Cys Ser Leu Phe Phe Leu Leu Trp  $1 \hspace{1.5cm} 5 \hspace{1.5cm} 10 \hspace{1.5cm} 15$ Val Leu Ala Glu Pro Ala Glu Asn Ser Asp Phe Tyr Leu Pro Gly Asp 20 25 30Tyr Leu Leu Gly Gly Leu Phe Ser Leu His Ala Asn Met Lys Gly Ile Val His Leu Asn Phe Leu Gln Val Pro Met Cys Lys Glu Tyr Glu Val Lys Val Ile Gly Tyr Asn Leu Met Gln Ala Met Arg Phe Ala Val Glu 65 70 75 80Glu Ile Asn Asn Asp Ser Ser Leu Leu Pro Gly Val Leu Leu Gly Tyr Glu Ile Val Asp Val Cys Tyr Ile Ser Asn Asn Val Gln Pro Val Leu Tyr Phe Leu Ala His Glu Asp Asn Leu Leu Pro Ile Gln Glu Asp Tyr Ser Asn Tyr Ile Ser Arg Val Val Ala Val Ile Gly Pro Asp Asn Ser Glu Ser Val Met Thr Val Ala Asn Phe Leu Ser Leu Phe Leu Leu Pro Gln Ile Thr Tyr Ser Ala Ile Ser Asp Glu Leu Arg Asp Lys Val Arg 170

Phe Pro Ala Leu Leu Arg Thr Thr Pro Ser Ala Asp His His Val Glu Ala Met Val Gln Leu Met Leu His Phe Arg Trp Asn Trp Ile Ile Val Leu Val Ser Ser Asp Thr Tyr Gly Arg Asp Asn Gly Gln Leu Leu Gly Glu Arg Val Ala Arg Arg Asp Ile Cys Ile Ala Phe Gln Glu Thr Leu Pro Thr Leu Gln Pro Asn Gln Asn Met Thr Ser Glu Glu Arg Gln Arg Leu Val Thr Ile Val Asp Lys Leu Gln Gln Ser Thr Ala Arg Val Val 265 Val Val Phe Ser Pro Asp Leu Thr Leu Tyr His Phe Phe Asn Glu Val Leu Arg Gln Asn Phe Thr Gly Ala Val Trp Ile Ala Ser Glu Ser Trp Ala Ile Asp Pro Val Leu His Asn Leu Thr Glu Leu Gly His Leu Gly Thr Phe Leu Gly Ile Thr Ile Gln Ser Val Pro Ile Pro Gly Phe Ser Glu Phe Arg Glu Trp Gly Pro Gln Ala Gly Pro Pro Pro Leu Ser Arg Thr Ser Gln Ser Tyr Thr Cys Asn Gln Glu Cys Asp Asn Cys Leu Asn Ala Thr Leu Ser Phe Asn Thr Ile Leu Arg Leu Ser Gly Glu Arg Val Val Tyr Ser Val Tyr Ser Ala Val Tyr Ala Val Ala His Ala Leu His Ser Leu Leu Gly Cys Asp Lys Ser Thr Cys Thr Lys Arg Val Val Tyr Pro Trp Gln Leu Leu Glu Glu Ile Trp Lys Val Asn Phe Thr Leu Leu Asp His Gln Ile Phe Phe Asp Pro Gln Gly Asp Val Ala Leu His Leu Glu Ile Val Gln Trp Gln Trp Asp Arg Ser Gln Asn Pro Phe Gln Ser

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450 455 460

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Asn	Pro	Asn	Tyr	Arg 725	Asn	Ser	Leu	Leu	Phe 730	Asn	Thr	Ser	Leu	Asp 735	Leu

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